User Manual for

TASSEL

Trait Analysis by aSSociation, Evolution and Linkage

Version 5.0

The Buckler Lab at Cornell University

(August 17, 2014)



www.maizegenetics.net/tassel

Disclaimer: While the Buckler Lab at Cornell University has performed extensive testing and results are, in general, reliable, correct or appropriate. Results are not guaranteed for any specific set of data. It is strongly recommended that users validate TASSEL results with other software.

Further help: Additional help is available beyond this document. Users are welcome to report bugs, request new features through the TASSEL website. Questions are also welcome to our current team members. For more quick and precise answers, please address your questions to the most pertinent person:

Tassel User Group (recommended)	http://groups.google.com/group/tassel tassel@googlegroups.com
General Information	Ed Buckler (Project leader)
Data Import, Pipeline	Terry Casstevens
Statistical Analysis	Peter Bradbury
	pjb39@cornell.edu

Contributors: Ed Buckler, Terry Casstevens, Peter Bradbury, Zhiwu Zhang, Dallas Kroon, Jeff Glaubitz, Kelly Swarts, Jason Wallace, Fei Lu, Alberto Romero, Cinta Romay, Eli Rodgers-Melnick, Alexander Lipka, Sara Miller, James Harriman, Yogesh Ramdoss, Michael Oak, Karin Holmberg, Natalie Stevens, and Yang Zhang.

Citations:

Overall Package:

Bradbury PJ, Zhang Z, Kroon DE, Casstevens TM, Ramdoss Y, Buckler ES. (2007) <u>TASSEL: Software for</u> association mapping of complex traits in diverse samples. Bioinformatics 23:2633-2635.

Genotyping by Sequencing:

Glaubitz JC, Casstevens TM, Lu F, Harriman J, Elshire RJ, Sun Q, Buckler ES. (2014) <u>TASSEL-GBS: A High</u> <u>Capacity Genotyping by Sequencing Analysis Pipeline</u>. *PLoS ONE* **9**(2): e90346

Mixed Model GWAS:

Zhang Z, Ersoz E, Lai C-Q, Todhunter RJ, Tiwari HK, Gore MA, Bradbury PJ, Yu J, Arnett DK, Ordovas JM, Buckler ES. (2010) <u>Mixed linear model approach adapted for genome-wide association studies</u>. *Nature Genetics* **42**:355-360. The TASSEL project is supported by the National Science Foundation and the USDA-ARS.





Reference Links:

Main Web Site: <u>http://www.maizegenetics.net/tassel</u> Open source code: <u>https://bitbucket.org/tasseladmin/tassel-5-source</u> Wiki: <u>https://bitbucket.org/tasseladmin/tassel-5-source/wiki</u>

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Introduction

While TASSEL has changed considerably since its initial public release in 2001, its primary function continues to be providing tools to investigate the relationship between phenotypes and genotypes¹. TASSEL has functionality for association study, evaluating evolutionary relationships, analysis of linkage disequilibrium, principal component analysis, cluster analysis, missing data imputation and data visualization. TASSEL development has been led by a group focused on maize genetics and genomics, and for these reasons that software has design and computational optimizations that account for the biology found in many plants and breeding situations. Compared to human genetics, many crops are highly diverse both at the nucleotide level and structural variations (10-50X greater than humans), inbreeding is common, large families are common, and whole genome prediction is being applied daily to real world problems. These biological differences lead to some different optimizations that are of use to many biological systems outside of crops.

One of the design elements driving TASSEL development has been the need to analyze ever larger sets of data². TASSEL5 has at its heart lots of design optimizations for big data, including:

- Bit level encoding of nucleotides so genetic distance and linkage disequilibrium estimates can be made very quickly (20-50X speed increases).
- Extensive use the HDF5 file format, which has been developed as a robust element of many climate modelers for matrix style data
- Tools for extracting and calling SNPs from extensive Genotyping-by-Sequencing data (tested for 60,000 samples by over 2.5 million SNPs and 96 million sequence alleles).
- Projection and imputation procedures that are optimized for the large families in crops. Some of these optimizations permit memory and computational improvements of >100,000 fold.
- Mixed models based on DNA relationships have come to dominate GWP (Meuwissen et al 2001) and GWAS (Yu et al 2006), yet these models can be slow to solve. TASSEL has been a test bed and implements some of the most best optimizations, such as EMMA (Kang at al 2008), plus approaches optimize variance components once P3D (Zhang et al 2010) and EMMAX (Kang et al 2010). Compression algorithms are also available (Zhang et al 2010). When used correctly, these optimizations make powerful GWAS computationally possible.
- The code is being continually optimized for larger numbers of cores and clusters. For example, we generally run imputation on 64-core machines. And while Java provides some excellent is interoperability between systems, its code is about 2-fold slower than optimized C libraries, and 10-fold slower than GPU processing for some problems. TASSEL5 is building out connection layers directly to native code, when these efficiencies are need.

TASSEL was designed for a wide range of users, including those not expert in statistical genetics or computer science. A GWAS using the mixed linear model method to incorporate information about population structure⁶⁻⁸ and cryptic relationships⁹ can be performed by in a few steps by "clicking" on the proper choices using a graphic interface. All the processes necessary for the analysis are performed automatically, including importing phenotypic and genotype data, imputing missing data (phenotype or genotype), filtering markers on minor allele frequency, generating principal components and a kinship matrix to represent population structure and cryptic relationships, optimizing compression level and performing GWAS.

The command-line version of TASSEL, called the Pipeline, provides users the ability to program tasks using a script instead of the graphic user interface (GUI). This feature allows researchers to define tasks using a few lines of code and provides the ability to use TASSEL as part of an analysis pipeline or to perform simulation studies.

We are also building a larger community of scientist developers that are adding functionality to this platform and working together to improve the system. So throughout this user manual you will see how to do most things three different ways - with the GUI, with the pipeline, and with the API (application programming interface).

TASSEL is written in Java, thereby enabling its use with virtually any operating system. It can be installed using Java Web Start technology by simply clicking on a link at <u>www.maizegenetics.net/tassel</u>. A stand-alone version of TASSEL can also be downloaded to use in pipeline mode or in any situation where the user wishes to start the software from a command line.

Getting Started

A quick way to get started using TASSEL is to load the tutorial data and try performing analyses. However, because some of the necessary steps may not be intuitive, we recommend that new users follow the tutorial at end of this manual. The objective of this section is to provide information necessary to install and start TASSEL software and to provide a brief overview of the interface.



class net.maizegenetics.dna.snp.CoreGenotypeTable

1.1 Executing TASSEL

http://www.maizegenetics.net/tassel/docs/ExecutingTassel.pdf

1.2 Open Source Code

Open source code for TASSEL is available at: <u>https://bitbucket.org/tasseladmin/tassel-5-source</u>. The package uses a number of other libraries that are included in the TASSEL distribution. These include a modified version of the PAL library (<u>http://www.cebl.auckland.ac.nz/pal-project/</u>), the COLT library (<u>http://dsd.lbl.gov/~hoschek/colt/</u>), jFreeChart (<u>http://www.jfree.org/jfreechart/</u>), Guava (Google Core Libraries) (<u>https://code.google.com/p/guava-libraries</u>), JUnit (<u>http://junit.org</u>), Archaeopteryx (<u>https://sites.google.com/site/cmzmasek/home/software/archaeopteryx</u>), and BioJava (<u>http://www.biojava.org</u>).

1.3 Software Development Tools

jProfiler (http://www.ej-technologies.com/products/jprofiler/overview.html) install4j (http://www.ej-technologies.com/products/install4j/overview.html) NetBeans IDE (https://netbeans.org) Eclipse (http://www.eclipse.org) IntelliJ (http://www.jetbrains.com/idea) Structure101 (http://structure101.com) TeamViewer (http://structure101.com) Bitbucket (https://bitbucket.org) sourceforge (http://sourceforge.net) JIRA (https://www.atlassian.com/software/jira) Tower (http://www.git-tower.com)

1.4 Graphical Interface

TASSEL is organized into five main panels. 1) At the top menus control functions. 2) The Data Tree at the top left organizes data sets and results. Data set(s) displayed in the Data Tree must first be selected before a desired function or analysis can be performed. To select multiple data sets, press the CTRL (or Command for Mac) key while selecting the data sets. 3) The Report Panel is located below the Data Tree. It displays information about a selected data set from the Data Tree, such as the type of data and how it was created. 4) The Progress Monitoring Panel below the Report Panel shows the progress of running tasks and has buttons that can cancel tasks. 5) The Main Panel occupies the right side of the viewing area, and displays the content of the selected data set from the Data Tree.

1.5 Pipeline (Command Line Interface)

http://www.maizegenetics.net/tassel/docs/TasselPipelineCLI.pdf

1.6 GBS Pipeline

http://www.maizegenetics.net/tassel/docs/TasselPipelineGBS.pdf

2 File Menu

The data tree can be saved in a binary format.

2.1.1 Save Data Tree

This feature allows you to save the entire contents of the Data Tree panel to a default location. This is helpful when the user does not wish to recreate a Data Tree panel that is already well populated with information the next time they initializes the program. To save a Data Tree, select **File > Save Data Tree**.

2.1.2 Open Data Tree

To restore a Data Tree that was saved previously saved, select File > Open Data Tree.

2.1.3 Save Data Tree As...

To save the Data Tree to a specific location or to give it a specific name, select File > Save Data Tree As....

2.1.4 Open Data Tree...

To restore a Data Tree from a specific location, select File > Open Data Tree...

NOTE: The information outlined above for saving a Data Tree is applicable to files that are, in general, version specific. When a new version of TASSEL is released, a data tree saved with a previous version might not load to the version. For longer term storage, the best practice is to save individual data sets rather than the entire data tree.

2.1.5 Set Preferences

Currently there is only one preference. That is whether to retain "rare" alleles. This is irrelevant for nucleotide data (A, C, G, T, -, +, N) because at that number of states, there is no data lost. Potentially with other types of data, it could exceed the 14 max (per site) number of allele states. If you "Retain Rare Alleles", the lower frequency allele values will be consolidated into a rare (Z) state. Otherwise, those lower frequency alleles are changed to unknown (N).

⊖ ○ ⊖ Preferences
Alignment Preferences
🗹 Retain Rare Alleles
OK Cancel

3 Data Menu

The Data Menu has options to import and export data sets, as well as, other data manipulate functions.

3.1 Load

Load provides options to import files for genotypes, phenotypes, populations structure, and kinship matrices, etc.

The tutorial data can be downloaded from the TASSEL website at this link:

http://www.maizegenetics.net/tassel/docs/TASSELTutorialData3.zip.

To use the data, the zip file must be uncompressed and saved on your local machine. These tutorial files will load correctly with the "Make Best Guess" option. Multiple files can be imported simultaneously by highlighting them first (holding Shift or Control key while clicking) and then clicking the Open button.



3.1.1 Hapmap

Hapmap is a text based file format for storing sequence data. All the information for a series of SNPs as well as the germplasm lines are stored in one file. The first row contains the header labels, and each additional row contains all the information associated with a single SNP. The first 11 columns describe attributes of the SNP, while the following columns describe the SNP value for a single germplasm line. The first 12 columns of the first

row should look like this, where "Line 1" is the beginning of germplasm line names.

rs#	alleles	chrom	pos	strand	assembly#	center	protLSID	assayLSID	panelLSID	QCcode	Line 1
-----	---------	-------	-----	--------	-----------	--------	----------	-----------	-----------	--------	--------

While all 11 header columns are required, not all 11 of the columns need to be filled in for TASSEL to correctly interpret the data. The only required fields are "chrom", Chromosome name, and "pos", Position. In the example below, genotype values are represented by 2 characters (i.e. AA). Note that you can record those as single character values (see "Nucleotide Codes" in the Appendix).

For TASSEL to correctly read Hapmap data, the data must be in order of position within each chromosome, and the file should be TAB delimited (example below is in Excel only for easy viewing). If some of the data is missing the correct number of TABs must still be present, so that TASSEL can properly assign data to columns.

rs#	alleles	chrom	pos	strand	assembly#	center	protLSID	assayLSID	panel	QCcode	33-16	38-11	4226	4722	A188
PZB00859.1	A/C	1	157104	+	AGPv1	Panzea	NA	NA	maize282	NA	CC	CC	CC	CC	AA
PZA01271.1	C/G	1	1947984	+	AGPv1	Panzea	NA	NA	maize282	NA	CC	GG	CC	GG	CC
PZA03613.2	G/T	1	2914066	+	AGPv1	Panzea	NA	NA	maize282	NA	GG	GG	GG	GG	GG
PZA03613.1	A/T	1	2914171	+	AGPv1	Panzea	NA	NA	maize282	NA	TT	TT	TT	TT	TT
PZA03614.2	A/G	1	2915078	+	AGPv1	Panzea	NA	NA	maize282	NA	GG	GG	GG	GG	GG
PZA03614.1	A/T	1	2915242	+	AGPv1	Panzea	NA	NA	maize282	NA	TT	TT	TT	TT	TT
PZA00258.3	C/G	1	2973508	+	AGPv1	Panzea	NA	NA	maize282	NA	GG	CC	CC	CG	CC
PZA02962.13	A/T	1	3205252	+	AGPv1	Panzea	NA	NA	maize282	NA	π	π	TT	π	π
PZA02962.14	C/G	1	3205262	+	AGPv1	Panzea	NA	NA	maize282	NA	CC	CC	CC	CC	CC
PZA00599.25	C/T	1	3206090	+	AGPv1	Panzea	NA	NA	maize282	NA	CC	π	CC	π	π
PZA02129.1	C/T	1	3706018	+	AGPv1	Panzea	NA	NA	maize282	NA	TT	CC	CC	CC	CC
PZA00393.1	C/T	1	4175293	+	AGPv1	Panzea	NA	NA	maize282	NA	TT	π	TT	CC	TT
PZA02869.8	C/T	1	4429897	+	AGPv1	Panzea	NA	NA	maize282	NA	CC	π	CC	NN	CC
PZA02869.4	C/G	1	4429927	+	AGPv1	Panzea	NA	NA	maize282	NA	CC	CC	CC	NN	GG
PZA02869.2	C/T	1	4430055	+	AGPv1	Panzea	NA	NA	maize282	NA	NN	π	TT	CC	π
PZA02032.1	A/T	1	4490461	+	AGPv1	Panzea	NA	NA	maize282	NA	AA	TT	AA	AA	AA
zagl1.5	A/T	1	4835434	+	AGPv1	Panzea	NA	NA	maize282	NA	AA	NN	AA	AA	AA
zagl1.2	A/C	1	4835558	+	AGPv1	Panzea	NA	NA	maize282	NA	CC	CC	CC	CC	CC
zagl1.6	C/T	1	4835658	+	AGPv1	Panzea	NA	NA	maize282	NA	TT	TT	TT	TT	TT
PZD00081.2	C/T	1	4836542	+	AGPv1	Panzea	NA	NA	maize282	NA	CC	CC	CC	CC	CC
zagl1.1	A/C	1	4912526	+	AGPv1	Panzea	NA	NA	maize282	NA	AA	AA	AA	AA	AA
PZB00919.1	A/C	1	5353319	+	AGPv1	Panzea	NA	NA	maize282	NA	CC	CC	CC	CC	AA
PZB00919.2	G/T	1	5353655	+	AGPv1	Panzea	NA	NA	maize282	NA	GG	GG	GG	GG	GG

3.1.2 HDF5 (Hierarchical Data Format version 5)

http://www.hdfgroup.org/HDF5

3.1.3 VCF (Variant Call Format)

http://www.1000genomes.org/wiki/analysis/variant-call-format/vcf-variant-call-format-version-42

3.1.4 Plink

Plink is a whole genome association analysis tool set, which comes with its own text based data format. The data is stored in a set of two files, a .map file and a .ped file.

The .ped file contains all the SNP values and has six mandatory header columns for Family ID, Individual ID,

Paternal ID, Maternal ID, Sex and Phenotype. TASSEL only requires that the Individual ID field be filled in. Each row of the .ped file describes a single germplasm line. Notice in Plink, an unknown character is represented with a '0'. However in TASSEL an unknown character is represented with a 'N', and '0' is used to represent heterozygous indel. TASSEL will automatically convert between the '0' and the 'N'. Any exported Plink files will represent the heterozygous indel with a '+' (insertion) and a '-' (deletion).

The .map file describes all the SNPs in the associated .ped file, where each row provides information on one SNP. The .map file must contain exactly four columns: Chromosome, rs#, Genetic distance and Position. TASSEL does not require the Genetic distance field to be filled in.

Both files should be TAB delimited.

For a more detailed description on the data format, please visit the Plink basic usage and data formats webpage: (http://pngu.mgh.harvard.edu/~purcell/plink/data.shtml).

3.1.5 Projection Alignment

3.1.6 Phylip

Details on Phylip format are described at the following website: <u>http://evolution.genetics.washington.edu/phylip/doc/sequence.html</u>

3.1.7 FASTA

3.1.8 Numerical Data

This type of format is used for trait and covariate data such as population structure. Similar to sequence alignment genotype data, numerical data also consists of two parts: a header that defines data structure and a body containing the main data. Tabs should be used as delimiters. However, any white space character such as blank will be treated as a delimiter as well. As a result, embedded blanks in names will cause data to be imported incorrectly. We suggest representing missing values using "NA", or "NaN". However, any text value (e.g. "?") will be interpreted as missing data. There are several formats for numerical data to fit the requirement for modeling. Trait data (dependent variables) can be imported by starting the first line with "<Trait>" and following that with the trait names. Additional classifiers may also be included in subsequent header rows by starting the row with "<Header name=xxx>" followed by a name for each column of data. For instance, to define environments, start the second header row with "<Header name=env>".

Comment lines may be inserted at the beginning of the file. Comment line begins with the character "#".

3.1.8.1 Trait format

This format does not require users to provide information on number of rows and columns. The file starts with the key word <Trait> followed by names of columns. The column for line should not be labeled.

Example 1, simple list of trait values:

<Trait> EarHT dpoll EarDia 811 59.5 NA NA 33-16 64.75 64.5 NA 38-11 92.25 68.5 37.897 4226 65.5 59.5 32.21933 4722 81.13 71.5 32.421 A188 27.5 62 31.419 ...

Example 2, traits data collected in multiple environments:

<Trait> EarHT PlantHT EarHT PlantHt
<Header name=env> Loc1 Loc1 Loc2 Loc2
811 59.5 NA NA NA
33-16 64.75 121.5 NA NA
38-11 92.25 153.8 37.897 83.4
4226 65.5 130.1 32.21933 82.1
4722 81.13 165.7 32.421 90.1
A188 27.5 110.2 31.419 79.6
...

3.1.8.2 Covariate Format

Covariate data uses the same format as trait data except that the first line must be "<Covariate>". This line tells TASSEL that the variables in this file will be used as covariates not as dependent variables. This is the format to use for population structure covariates.

```
<Covariate>
<Trait> Q1 Q2 Q3
33-16 0.014 0.972 0.014
38-11 0.003 0.993 0.004
4226 0.071 0.917 0.012
4722 0.035 0.854 0.111
A188 0.013 0.982 0.005
...
```

3.1.8.3 Marker Values as Numerical Co-variates

In some cases, a user may wish to have marker values treated as numerical co-variates. If the first line of the file is "<Numeric>", then the data will be imported as numeric data but used as marker data in GLM and MLM.

```
<Numeric>
<Marker> m1 m2 m3 m4 m5
33-16 0 1 1 0 0
38-11 0 0 1 0.3 0
4226 0 1 1 0.5 0
```

3.1.9 Square Numerical Matrix

Kinship can be calculated externally from pedigrees by using SAS Proc Inbreeding¹⁸ or from markers by using one of several available software packages. The following format is provided to import the resulting kinship estimates:

If n represents the number of taxa, the format for kinship files is as follows:

n Taxa1Name r11 r12 ... r1n Taxa2Name r21 r22 ... r2n ... TaxanName rn1 rn2 ... rnn

Here rij (i, j=1,2, ..., n) is the element in the kinship matrix located at row i and column j.

Missing values are not allowed for kinship matrix.

Important note: The current format is different from the format used in TASSEL version 2.0 or lower.

3.1.10 Table Report

Data can be imported as tab delimited text files. The first row of the file will be interpreted as column labels and the remaining rows as rows in the table.

3.1.11 TOPM (Tags on Physical Map)

3.2 Export

Options are provided to export sequence data: Hapmap, Plink, Phylip (Sequential or Interleaved). Phenotypes and covariate data is exported as numerical trait data. Table Reports are exported as a tab delimited table. For numerical data, the function of Export is similar to the Table function in Results mode.

\odot	Write Ha	pmap
\bigcirc	Write HD	F5
\bigcirc	Write VC	F
\bigcirc	Write Plin	nk
\bigcirc	Write Phy	ylip (Sequential)
\bigcirc	Write Phy	ylip (Interleaved)
0	Nrite Tal	b Delimited

3.3 Sort Genotype File

TASSEL5 has strict requirements for the sites in a genotype file. Each site must be unique (as defined by its locus/chromosome, position, and name) and they must be in order in the file. Genotype files produced by other programs (and also earlier versions of TASSEL) often do not meet this second requirement and throw an error when TASSEL tries to load them. It can be difficult to recreate TASSEL's internal sort order by hand, so this plugin allows the user to sort an input genotype file according to TASSEL's rules and output it to a new file ready for further analysis. (This sort is not done automatically at load time because the computational cost for sorting large files can be very large. We feel it's better for users to know what they're getting into instead of being surprised by it.) There is currently only support for sorting Hapmap and VCF files.

To sort a genotype file from the GUI, just select Data -> Sort Genotype File and fill in the appropriate parameters in the popup dialog.

To sort a file from the command line, use the following command:

```
run_pipeline.pl -SortGenotypeTablePlugin -inputFile [filename] -outputFile
[filename] -fileType [Hapmap or VCF]
```

The -fileType flag is optional and is only needed if the input file's extension doesn't match a known file extension (".hmp.txt", ".vcf", etc.).

3.4 Transform

This suite of functions allows multiple data manipulation on genotype and phenotype (numerical) data. When a genotype data set is selected, the data are transformed to numbers. When a numerical data set is selected, mathematical transformation, data imputation and principal component analysis (PCA) can be performed. The Transform columns tags will be displayed in a Data dialog box with three tabs: Trans, Impute and PCA.

3.4.1 Genotype Numericalization

Two options are provided to transform genotype from character to numerical as shown in the following dialog box.



3.4.1.1 Collapse Non Major Alleles

This function assigns 1 to the major allele and 0 to any other alleles. The converted genotypes are saved in a new numerical data set.

3.4.1.2 Separate Alleles

This function assigns an indicator (1 for present and 0 for absent) for each allele. The converted genotypes are saved in a new numerical data set.

3.4.2 Transform and/or Standardize Data

The **Trans** dialog box is the default selection, as shown below. In the **Column** list, select the column(s) you wish to transform. Then select the type of transformation you wish to execute. Selecting the **Standardize** checkbox will transform data by subtracting the column mean from the value of the trait and then dividing by the column's standard deviation. Clicking on the **Create Data set** button will result in the placement of a dataset containing only the selected columns in the Data Tree.

<u>\$</u>	10 10.000	×
Column	Percent Missing Data	Trans Impute PCA
EarHT.null	0.66	Raise to Power 2
dpoll.null		
EarDia.null	12	O Take Log Base 10
		Standardize
	Create Dataset Close	

3.4.3 Impute Phenotype

The k-nearest-neighbor algorithm²⁰ is used to impute missing phenotype data. If data is missing for a taxon for one of the traits, the algorithm finds other taxa (neighbors) that are most like it for the non-missing traits. It uses the average of the neighbors to impute the missing data. Click on the **Impute** tab to display the following:

\$	11.145 10 10 10.000	×
Column	Percent Missing Data	Trans Impute PCA
EarHT.null	0.66	Manhatten Distance
dpoll.null	1.3	
EarDia.null	12	 Euclid Distance
		Our Unweighted Average
		Weighted Average
		Number of Neighbors (K): 3 🚔
		Min. Freq. of Row Data: 0.80
	Create Dataset Close	J

3.4.4 PCA

Principal component analysis (PCA) can only be performed on a numerical data set without missing values. Two methods are available: correlation or covariance. This determines whether a correlation or covariance matrix will be used as the basis for the analysis. The default, correlation, is a reasonable choice for genetic data. The number of PCA axes in the output data set can be controlled by selecting either of the minimum eigen value associated

with each axis, the minimum percent of the variance captured by an axis or the number of axes. The resulting axes will be sorted by the amount of variance each captures.

Т								
Column	Percent Missing Data	Trans Impute	PCA					
EarHT.NA	0.00							
dpoll.NA	0.00	Me	thod					
EarDia.NA	0.00	⊙ Co	rrelation					
		000	variance					
		0.	Jtput					
		💿 Eigenvalu	e≥ 0					
		🚫 Var Prop	% ≥ 0.33333					
		O Componer	nts = 3 🛟					
Create Dataset Close								

3.5 Synonymizer (Synonymize Taxa Names)

This button makes taxa names uniform to permit the joining of data sets.

The join functions that generate fused data sets work by matching taxa names. Consequently, if multiple names exist for a given taxon (an added suffix, alternative spellings, different naming conventions, etc.) then the two data sets will not join correctly. To help remedy this, the Synonymizer function allows the taxa names of one data set to replace similar taxa names in the second data set. It relies on an algorithm that calculates the degree of similarity between names, using the name from the first set which is most similar to that in the second data set.

When using the Synonymizer, keep in mind that order of selection matters. Always select the data set with the names you wish to use (the "real" name) *first*, and then, while holding down the CTRL key, click on the second data set with the taxa names you wish to change (the "synonym"). Then click on the **Synonymizer** button. A synonym data set will be placed on the Data Tree panel under **Synonyms**. Each name in the data set selected second is now listed in the **TaxaSynonym** column. Next to this column is a **TaxaRealName** column listing the highest scoring match derived from the "real" name data set. The **MatchScore** column gives an indication of the amount of similarity between the two names (where 0 is no similarity and 1.0 is identity).

TASSEL (Trait Analysis by aSSociation, Evolution, and Linkage) 3.0.36							
File Tools Help <u>G</u> DPC							
CARAGE CORRECTION OF Analysis Results Delete Wizard	M Sh	ow Memory				0%	
🖏 GDPC 🗐 Load 🏦 Export 🍸 Sites 🕨 Taxa 🍸 Traits 🗱 Ir	npute SNPs	?+5 Transform	p⇔q syno	onymizer	🗊 u Join	n Join	⊘⊘ Separate
Data 🔺	TaxaSyno	TaxaRealN	RefIDNum	MatchScore			
🚊 🤚 Sequence	T7111	1	-1	0.0			
d8_sequence	TZT 10	T7I 10	82	1.0	-		
mdp_genotype	B79	B73	12	0.5	-		
Polymorphisms	AB28A		-1	0.0	-		E
🖶 👘 Numerical	B77	B73	12	0.5	-		
mdp_population_structure	M37W	M37W	52	1.0	-		
- mdp_traits	B76	B73	12	0.5	-		
😑 👘 Matrix	KI3	KI3	47	1.0	-		
mdp_kinship	B75	B73	12	0.5	-		
• Tree	GT112	GT112	37	1.0	-		
+ Fusions	CI31A	KI3	47	0.33333333.			
Synonymizer	B73	B73	12	1.0	-		
d8_sequence Synonyms	VA59	A554	3	0.33333333.			
🐌 Result	CI44	KI44	49	0.66666666.			
	CML92	CML5	29	0.57142857.			
	CML91	CML5	29	0.57142857.			
	CI7	CI187-2	16	0.25	-		
Table Title: Taxa Synonym Table	CML5	CML5	29	1.0			
Number of columns: 4	SC55	SC55	78	1.0			
Number of rows: 301	OH7B	OH7B	72	1.0			
Number of elemental 1204	IL677A	IL677A	43	1.0			
Torre energiese izer	CI3A	KI3	47	0.4			
Taxa synonyms	CM37	M37W	52	0.66666666.			
Synonym Table	B68	B68	11	1.0			
301 unique matches	B115		-1	0.0			
73 unmatched:	B64	B68	11	0.5			
	H100		-1	0.0			
	DE811	KI11	45	0.28571428.			
	CH9	H99	38	0.5			
Load Cancel	CO255	K55	44	0.33333333.			
	NC372		-1	0.0			
	NC370		-1	0.0			
Synonymizer Cancel	U267Y	U267Y	85	1.0			
	B109		-1	0.0			
	WF9	WF9	90	1.0			-
Program Status							

Caution! Before the synonyms are applied, we strongly encourage the user to check the match score, especially for those taxa with low match scores. To do that, the user selects the synonym file and clicks the "Synonymizer" button. The incorrect matches, usually the ones with the lowest match scores, can be rejected at this point. Sorting on the match score column first makes this a fairly easy process.

In the event that some of the taxa are not interpreted correctly, matches can be modified manually. Select the taxa you wish to modify on the left side, and then choose a replacement taxa from the right side. Click the arrow button

to substitute the taxa. Taxa with no synonym can be identified by selecting then clicking "No Synonym". Click **OK** to save the changes.

Synor	nymizer						
TaxaSy	TaxaRe	RefIDNum	MatchSc		т	ZI 18	
TZI11		-1	0.0	i.	Т	ZI 10	
TZI 10	TZI 10	82	1.0		Т	ZI8	
B79	B73	12	0.5	-	ĸ	I11	
AB28A		-1	0.0		3	8-11	
B77	B73	12	0.5		G	T112	
M37W	M37W	52	1.0		N	V117HT	
B76	B73	12	0.5		I	137TN	
KI3	KI3	47	1.0		C	I 187-2	
B75	B73	12	0.5		N	VF9	
GT112	GT112	37	1.0		M	V64A	
CI31A	KI3	47	0.333333		< N	V182B	
B73	B73	12	1.0		N	V153R	
VA59	A554	3	0.333333		U	1267Y	
CI44	KT44	49	0.666666		T	X601	
CMI 92	CMI 5	29	0.571428		Т	232	
CMI 91	CML5	29	0.571428		S	G18	
CI7	CI 187-2	16	0.25		S	C55	
CML5	CMI 5	29	1.0		S	C213R	
SC55	SC55	78	1.0		S	A24	
OH7B	OH7B	72	1.0		Q	26199	
IL677A	IL677A	43	1.0	-	P	A91	
	No S	Synonym				Cape	Sort Alphabetically

Once it has been determined that the taxa names were matched correctly, the synonyms can be applied. With the synonyms selected, hold down the CTRL key while clicking on the second/synonym data set (the data set whose names you would like to change). Then once again click on the **Synonymizer** button to apply the new names to the data set.

3.6 Intersect Join

Command

```
./run_pipeline.pl -fork1 -h group1.hmp.txt -fork2 -h group2.hmp.txt
-combine3 -input1 -input2 -intersect -export
group1_group2_intersect.hmp.txt -runfork1 -runfork2 -runfork3
```

This joins multiple data sets by the intersection of their taxa. Taxa must be present in both data sets to be included. Select multiple data sets using the CTRL key in conjunction with mouse clicks, and then click on the intersection button to join the data sets. Because this function uses taxa names to join data sets, any variation in taxa names can

prevent proper joining. Taxa names can be made uniform by using the "Synonymizer".

3.7 Union Join

Command

```
./run_pipeline.pl -fork1 -h group1.hmp.txt -fork2 -h group2.hmp.txt
-combine3 -input1 -input2 -intersect -export group1_group2_union.hmp.txt
-runfork1 -runfork2 -runfork3
```

This joins multiple data sets by a union of their taxa. Missing data will be inserted if taxa are missing from one data set. Select multiple data sets using the CTRL key in conjunction with mouse clicks, and then click on the union button to join the data sets. Because this function uses taxa names to join data sets, any variation in taxa names can prevent proper joining. Taxa names can be made uniform by using the "Synonymizer".

3.8 Merge Genotype Tables

Command

```
./run_pipeline.pl -fork1 -h group1.hmp.txt -fork2 -h group2.hmp.txt
-combine3 -input1 -input2 -mergeAlignments -export
group1 group2 merge.hmp.txt -runfork1 -runfork2 -runfork3
```

This is the most complex merge function, and can be considered as a union join across both sites and taxa. (The actual -union join only works across taxa.) The resulting genotype table will contain all unique sites and all unique taxa from across the input datasets. If a specific site-taxon combination isn't present in any input dataset, the value is set to missing. If a specific site-taxon combination is present in more than one input file, the output will contain the last value processed. (That is, later values overwrite earlier values even if they conflict. There are plans to change this, but they have not been implemented yet.)

Notes

- This maps to "Data -> Merge Genotype Tables" Menu on GUI.
- Error if duplicate site names in same file. (same as with other file loadings)
- Undefined taxa / site allele values are set to UNKNOWN.
- Duplicate taxa / site set to last Alignment processed.
- Sites are identified by Locus (chromosome), Physical Position, and Site Name

3.9 Separate

This separates the selected data set into it's components. For example, a genotype table would be separated into individual chromosomes.

3.10 Homozygous Genotype

This changes all heterozygous values to unknown (N).

4 Impute Menu

4.1 Genotypic Imputation

TASSEL5 contains two methods for imputing missing genotype information, one is a generalized approach suitable for all types of populations but optimized for those with higher inbreeding coefficients (FILLIN) and the other is specifically optimized for finding recombination break points in full-sib families (FSFHap). More information on these two methods can be found at:

Swarts et al. FSFHap (Full-Sib Family Haplotype Imputation) and FILLIN (Fast, Inbred Line Library ImputatioN) optimize genotypic imputation for low-coverage, next-generation sequence data in crop plants, Plant Genome, *in review*.

FSFHap (Full-Sib Family Haplotype Imputation):

FSFHap imputes missing genotypes and corrects genotyping errors for inbred individuals in full-sib families. It is very useful for calling haplotypes in low-coverage GBS data. The individuals must be at least partially inbred because the method relies on finding inbred segments to identify haplotypes. It does not use the parent genotypes directly, but including the parents may be useful for interpreting the results. The algorithms used for imputation analyze one chromosome and family at a time. As a result, a pedigree file must be supplied that indicates which entries belong to which family. Also, input genotypes must contain data for only a single chromosome. If the genotype file contains multiple chromosomes, the chromosomes can be separated using the TASSEL -separate command.

Pedigree File Format:

The only file format specific to FSFHap is the pedigree file. The taxa names must exactly match names in the genotype data. If the genotype data contains taxa not included in the pedigree file, only individuals listed in the pedigree file will be analyzed. The input genotypes can be in any of the formats accepted by TASSEL. The pedigree file must contain the names of the individual taxa to be analyzed, the family to which each belongs, the parents, the parent contributions, and the average inbreeding coefficient. The first row in the file must be column headers. The values in the columns should be tab-delimited and are expected to be in the following order: family, taxon, parent1, parent2, parent1Contribution, parent2Contribution, F. The F value is not required but all other columns are.

Example:

family	taxonName	parentl	parent2	contribution1	contribution2	F
fam1	t0001	par1	par2	0.5	0.5	.92
fam1	t0002	parl	par2	0.5	0.5	.92
fam2	t0201	par1	par3	0.5	0.5	.92

fam2	t0202	par1	par3	0.5	0.5	.92
fam2	t0203	par1	par3	0.5	0.5	.92

The values for contribution1, contribution2, and F are family means. Those values are read from the first line for a family only and then applied to the entire family.

Using the command line for FSFHap:

FSFHap consists of three TASSEL plugins, CallParentAllelesPlugin, ViterbiAlgorithmPlugin, and WritePopulationAlignmentPlugin, which are called sequentially. A typical command for running FSFHap is as follows (replace items in > with actual parameter values) for a genotype containing a single chromosome:

run_pipeline.pl -h <genotypeFilename> -CallParentAllelesPlugin -p <pedigreeFilename> -m 0.9 -r 0.5 -logfile <logFilename> -endPlugin -ViterbiAlgorithmPlugin -g true -endPlugin -WritePopulationAlignmentPlugin -f <outputFilename> -m false -o parents -endPlugin

For a genotype file containing multiple chromosomes:

run_pipeline.pl -h <genotypeFilename> -separate -CallParentAllelesPlugin -p <pedigreeFilename> -m 0.9 -r 0.5 -logfile <logFilename> -endPlugin -ViterbiAlgorithmPlugin -g true -endPlugin -WritePopulationAlignmentPlugin -f <outputFilename> -m false -o parents -endPlugin

Options for CallParentAllelesPlugin:

- F ···· J · ··· · ·	
Options taking a para	meter value specified by Value = []:
-p or -pedigrees	the pedigree file. Value = [filename]
-w or -windowSize	the number of SNPs to examine for each LD cluster.
	Value = [integer] (default = 50)
-r or -minR	minimum R used to filter SNPs on LD
	Value = [number between 0 and 1]. (default = 0.2 , use 0 for no ld filter)
-m or -maxMissing	maximum proportion of missing data allowed for a SNP
	Value = [number between 0 and 1]. (default = 0.9)
-f or -minMaf	minimum minor allele frequency used to filter SNPs. If negative, filters on
	expected segregation ratio from parental contribution.
	Value = [number between 1 and -1]. (default = -1)
-b or -bc1	use BC1 specific filter. Value = [true or false] (default = true)
-n or -bcn	use multiple backcross specific filter. Value = [true or false] (default = false)
-logfile	the name of a file to which all logged messages will be printed. Value = [filename].

Options not taking a parameter value:

-cluster	use the cluster algorithm. minMaf defaults to 0.05.
-subpops	filter sites for heterozygosity in subpopulations.
-nohets	delete het calls from original data before imputing.
-windowld	use the window ld algorithm for finding parent haplotypes

The "-cluster", "-subpops", "-nohets", and "-windowld" options do not take parameters but only act as flags that include certain features in the analysis. Of those, cluster and windowld are the most useful. When the -cluster option is used, a different algorithm is used that does a better job of handling residual heterozygosity in the

parents. However, it does not perform well for partially inbred RILs that have only been self-pollinated for one or two generations. If the RILs being imputed are F2's or F3's, the "-cluster" option should not be used. The "-subpops" option should only be used when imputing families of the NAM population developed by the Maize Diversity Project. The "-nohets" option was included to test whether or not erroneous het calls result in too many hets being imputed. It appears to have only a small effect on the outcome. The -windowld algorithm handles F2 and later populations effectively, but can have problems when parents have some residual heterozygosity.

It is recommended that the -logfile option be used. The output can be used to identify and diagnose possible problems. The "-bcn true" should be used for populations with two or more backcrosses. However, using the "-bc1" option is not necessary as the default behavior is usually best.

Options for ViterbiAlgorithmPlugin:

-g or -fillgaps	if true then missing values flanked by SNPs from the same parent will be imputed
	to that parent, false otherwise. Value = [true or false] (default = true)
-h or -phet	expected frequency of heterozygous loci. Used only if the inbreeding coefficient is
	not specified in the pedigree file. Value = [number between 0 and 1] (default = 0.07)

Options for WritePopulationAlignmentsPlugin:

The base file name for the ouputhmp.txt will be appended. Value = [filename]
if true then families are merged into a single file, if false then each family is output
to a separate file. Value = [true or false] (default = false)
if value = parents then output parent calls, if value = nucleotides then output nucleotides,
if value = both then output both in separate files (default = both)
if true output is AA/CC/AC, if false output is A/C/M.
Value = [true or false] (default = false)
the minimum coverage for a monomorphic snp to be included in the nucleotide output.
Value = [number between 0 and 1] (default = 0.1)
the maximum minor allele frequency used to call monomorphic snps (default = 0.01)

For individual families, only polymorphic SNPs are imputed. When merge = false, only those SNPs appear in the output. When merge = true, SNPs that are polymorphic in any family will be written to output. For any site, if SNP coverage is high enough in a family to determine with confidence that it is monomorphic for that family, then all individuals in that family will be imputed to the monomorphic value at that site. The -minCoverage and -maxMono options are used to determine thresholds for determining whether a site will be called monomorphic in a family. If either of the options is set to a value of NaN, then missing values at monomorphic sites will not be imputed.

FILLIN (Fast, Inbred Line Library ImputatioN): The generalized approach

FILLIN imputes missing genotypes in two steps, 1) haplotype generation (FILLINFindHaplotypesPlugin) and 2) imputation of the resulting haplotypes back onto the target samples (FILLINImputationPlugin).

Haplotypes are generated by collapsing low coverage but inbred segments that share identity by state to an optionally user-supplied threshold value by site window (default: 8k); this is performed by the first plugin, FILLINFindHaplotypesPlugin. Because short IBD segments may be replicated widely within a species, even between diverse individuals, we recommend supplying all the information available within a species for this step.

The second plugin, FILLINImputationPlugin, uses these haplotypes to impute missing genotypes in target individuals. It does so in multiple steps, first looking for haplotypes that match the minor alleles to a threshold within the whole site window (1a in schematic below) and, if this fails, looks for two haplotypes to explain the site window and, assuming this represents a recombination break point between two inbred haplotypes, uses a Viterbi HMM algorithm to model the recombination breakpoints (2a). If two haplotypes cannot be found to explain the whole site window, the algorithm next searches for haplotypes to explain a smaller focus window within the site window centered on 64 sites at a time and searching to the right and left until enough informative minor alleles are found. It does this by first looking for one haplotype to a threshold (2a), then two modeling a recombination break between inbred segments (2b), then finally, to a higher threshold, looks for two haplotypes and models the 64 focus site window as heterozygous, combining the two haplotypes together. The thresholds for 2a-c are also set differently based on whether the whole sequence of the target taxon is above or below a user supplied heterozygosity threshold. For taxon considered outbred (above the threshold), 2b the Viterbi option is never used because it is more likely in an outbred taxon that if two haplotypes explain a segment it is heterozygous for those two haplotypes. If the algorithm cannot find haplotypes to satisfy any of these threshold requirements, the segment will not be imputed. The thresholds for the focus block imputation are set based on the mxInbErr and mxHybErr values entered (or defaults):

	Below mxHet (inbred)	Above mxHet (outbred)		
2a	3/10*mxInbErr	1/10*mxInbErr		
2b	⅓*mxHybErr	0		
2c	mxInbErr	mxInbErr		

FILLINFindHaplotypesPlugin

FILLINImputationPlugin



Running FILLIN:

FILLIN consists of two TASSEL plugins, FILLINFindHaplotypesPlugin and FILLINImputationPlugin, which are called sequentially. If you would like to mask your data and calculate accuracy, use the -accuracy flag for FILLINImputationPlugin. If imputing maize, a donor file of haplotypes from 40k+ taxa can be found on the Panzea website (<u>http://www.panzea.org/lit/data_sets.html</u>). FILLIN can be run either within the TASSEL GUI or through the command line. The options are the same for both.

A typical command sequence for running FILLIN through the command line is as follows (replace items in > with actual parameter values):

run_pipeline.pl -FILLINFindHaplotypesPlugin -hmp <genotypeFilename> -o <outDonorDir> run_pipeline.pl -FILLINImputationPlugin -hmp <genotypeFilename> -d <donorDir> -o <outFile.hmp.txt.gz>

To run FILLIN from the GUI go to Impute->FILLINFindHaplotypesPlugin or FILLINImputationPlugin

Options for FILLINFindHaplotypesPlugin:

-hmp <Target file> :

Input genotypes to generate haplotypes from. Usually best to use all available samples from a species. Accepts all file types supported by TASSEL5. (required)

-o <Donor dir/file basename> :

Output file directory name, or new directory path; Directory will be created, if doesn't exist. Outfiles will be placed in the directory and given the same name and appended with the substring '.gc#s#.hmp.txt' to

denote chromosome and section (required)

-mxDiv <Max divergence from founder> :

Maximum genetic divergence from founder haplotype to cluster sequences (Default: 0.01) -mxHet <Max heterozygosity of output haplotypes> :

Maximum heterozygosity of output haplotype. Heterozygosity results from clustering sequences that either have residual heterozygosity or clustering sequences that do not share all minor alleles. (Default: 0.01) -minSites <Min sites to cluster> :

The minimum number of sites present in two taxa to compare genetic distance to evaluate similarity for clustering (Default: 50)

-mxErr <Max combined error to impute two donors> :

The maximum genetic divergence allowable to cluster taxa (Default: 0.05)

-hapSize <Preferred haplotype size> :

Preferred haplotype block size in sites (minimum 64); will use the closest multiple of 64 at or below the supplied value (Default: 8192)

-minPres <Min sites to test match> :

Minimum number of present sites within input sequence to do the search (Default: 500)

-maxHap <Max haplotypes per segment> :

Maximum number of haplotypes per segment (Default: 3000)

-minTaxa <Min taxa to generate a haplotype> :

Minimum number of taxa to generate a haplotype (Default: 2)

-maxOutMiss <Max frequency missing per haplotype> :

Maximum frequency of missing data in the output haplotype (Default: 0.4)

-nV <true | false> :

Supress system out (Default: false)

-extOut <true | false> :

Details of taxa included in each haplotype to system out (Default: false)

Options for FILLINImputationPlugin:

-hmp <Target file> :

Input HapMap file of target genotypes to impute. Accepts all file types supported by TASSEL5 (required) -d <Donor Dir> :

Directory containing donor haplotype files from output of FILLINFindHaplotypesPlugin. All files with '.gc' in the filename will be read in, only those with matching sites are used (required)

-o <Output filename> :

Output file; hmp.txt.gz and .hmp.h5 accepted. (required)

-hapSize <Preferred haplotype size> :

Preferred haplotype block size in sites (use same as in FILLINFindHaplotypesPlugin) (Default: 8000) -hetThresh <Heterozygosity threshold> :

Threshold per taxon heterozygosity for treating taxon as heterozygous (no Viterbi, het thresholds). (Default: 0.01)

-mxInbErr <Max error to impute one donor> :

Maximum error rate for applying one haplotype to entire site window (Default: 0.01)

-mxHybErr <Max combined error to impute two donors> :

Maximum error rate for applying Viterbi with to haplotypes to entire site window (Default: 0.003) -mnTestSite <Min sites to test match> :

Minimum number of sites to test for IBS between haplotype and target in focus block (Default: 20) -minMnCnt <Min num of minor alleles to compare> :

Minimum number of informative minor alleles in the search window (or 10X major) (Default: 20) -mxDonH <Max donor hypotheses> :

Maximum number of donor hypotheses to be explored (Default: 20) -hybNN <true | false> :

If true, uses combination mode in focus block, else does not impute (Default: true) -ProjA <true | false> :

Create a projection alignment for high density markers (Default: false) -impDonor <true | false> :

Impute the donor file itself (Default: false)

-nV <true | false> :

Supress system out (Default: false)

Options for calculating accuracy

-accuracy <true | false> :

Masks input file before imputation and calculates accuracy based on masked genotypes (Default: false) -propSitesMask <Proportion of genotypes to mask if no depth> :

Proportion of genotypes to mask for accuracy calculation if depth not available (Default: 0.01) -depthMask <Depth of genotypes to mask> :

Depth of genotypes to mask for accuracy calculation if depth information available (Default: 9) -propDepthSitesMask <Proportion of depth genotypes to mask> :

Proportion of genotypes of given depth to mask for accuracy calculation if depth available (Default: 0.2)

5 Filter Menu

5.1 Sites

The genotype table can be filtered in several ways. For example, monomorphic sites can be eliminated, and regions of a sequence can be eliminated.

\varTheta 🔿 🖸 🖉 Filte	r Alignment	
	Filter Alignment	
Minimum Count:	10	out of 281 sequences
Minimum Frequency:	0.1	
Maximum Frequency:	1.0	
Position Type:	Position index	
Start Position:	0	
End Position:	3092 of 3092 sites	
Remove minor SNP states Generate haplotypes via sliding window		
Haplotype Length		
Step Length		
Filter Selec	t Chromosomes	Cancel

Minimum Count - the minimum number of taxa in which the site must have been scored to be included in the filtered data set (GAP or missing data do not count).

Minimum Frequency - the minimum frequency of the minority polymorphisms for the site to be included in the filtered data set.

Start Position, End Position – establishes the range of sites for filtering.

Extract Indels - if selected, indels are extracted from the alignment. If not selected, only point substitutions are extracted.

Remove minor SNP states – converts tertiary and rarer states to missing data ("?"), thereby forcing sites to have only two types of segregating sites at a locus. This may help remove sequencing errors.

Generate haplotypes via sliding window – creates haplotypes from an ordered set of SNPs.

Example Pipeline Command that removes SNPs with MAF (Minimum Allele Frequency) less than 5%

```
run_pipeline.pl -fork1 -h mdp_genotype.hmp.txt -filterAlign
-filterAlignMinFreq 0.05 -export filtered_genotype -runfork1
```

5.2 Site Names

First select the genotypic data from the data tree. The resulting dialog displays the site names associated with the selected data. By using either the CTRL or SHIFT key in conjunction with the mouse, the user can select or deselect site names. Once desired site names have been moved to the "Selected" window using the "Add ->" button, the "Capture Selected" or "Capture Unselected" buttons will create a new data set containing only the desired site names.

Using the search box...

- * is the wildcard.
- * is always implied at end of search string.
- Search string is case sensitive. For example: use [Aa]bc to match site names beginning with Abc or abc.
- PZ[AB] Will match anything starting with PZA or PZB.



5.3 Taxa Names

First select the genotypic, phenotypic, or population structure data from the data tree. The resulting dialog displays the taxa associated with the selected data. By using either the CTRL or SHIFT key in conjunction with the mouse, the user can select or deselect taxa. Once desired taxa have been moved to the "Selected" window using the "Add ->" button, the "Capture Selected" or "Capture Unselected" buttons will create a new data set containing only the desired taxa.

Using the search box...

- * is the wildcard.
- * is always implied at end of search string.
- Search string is case sensitive. For example: use [Aa]bc to match taxa beginning with Abc or abc.

Available		Selected	
A554	6	A556	
A556		A6	
A6		A619	
A619		A632	
A632	Add ->	A634	
A634			
A635			
A641	A		
A654	T		
A[56]			
Capture Selected	Capture Unselected	Remove	Cancel

A[56] Will match anything starting with A5 or A6

5.4 Taxa

$\Theta \cap \Theta$
Filter Taxa by Properties
Min Proportion of Sites Present 0.99
Min Heterozygous Proportion 0.0
Max Heterozygous Proportion 1.0
Ok Close

5.5 Traits

Clicking the "Traits" button on the "Data" toolbar launches the Trait Filter dialog. This dialog is used with numerical data sets to (1) change the trait type, (2) view, but not change whether the trait is discrete or continuous and (3) drop one or more traits from the data set. In addition, the dialog can be used to view the trait properties without changing them. If the "OK" button is clicked, a new data set is created that incorporates the changes, the original data set remains unchanged, and the dialog closes. If the "Cancel" button is clicked no data set is created, the original data set remains unchanged, and the dialog closes.

Allowable trait types are data, covariate, factor and marker. Generally, data and covariate traits will be continuous (not discrete) and factor will be discrete. Markers in a numerical data set will be continuous. Discrete valued

markers are better imported as genotypes and filtered using the "Sites" filter.

Clicking "Exclude All" unchecks the "Include" box for all traits. Clicking "Include All" checks the "Include" box for all traits. The "Exclude Selected" and "Include Selected" buttons do the same thing for traits that have been highlighted by selecting them with the mouse. Type can be changed for individual traits by selecting a value in the drop down box in the type column for that trait. Type can be changed for multiple traits by selecting those traits then clicking one of the "Change Selected Type to …" buttons.

Important: Once a numerical data set has been joined with genotypes, it can no longer be modified using the trait filter function.

9	Filter Traits / Modify Trait Properties				
Trait 01	Type	Discrete	Include		
Q2	covariate	Ö			
Q5	covariate				
	Exclude Selected	Include S	elected		
	Exclude All	Include A			
	Change Sel	ected Type to Dat	a		
	Change Selec	ted Type to Covari	iate		
	enunge seree				
	Change Sele	cted Type to Mark	er		
		Grand			
	OK	Cancel			

6 Analysis Menu

6.1 Diversity

This executes a basic diversity analysis.

Average pairwise divergence (π), segregating sites, and θ estimates (4Nµ) can be calculated, as well as sliding windows of diversity.

Diversity Surveys	×
Silent	
Noncoding	
Intron	Start Base 0
Synonymous	End Base 2465
Non-transcribed	
Noncoding Indels	
Transcribed	Sliding Window
Nonsynonymous	Step 100
Coding	Window 500
Coding Indels	WINDOW
Veral	
Indels	
Run	Close

To run a diversity analysis, click on a raw sequence alignment, and then select Analysis -> Diversity.

In the resulting Diversity Surveys dialog box, the various site classes available for analysis are listed on the left. If the sequence has no annotation, then only the "Overall" and "Indels" options will be active.

A sliding window of diversity can also be calculated across the region. To produce a sliding window, check the box next to "Sliding Window," and then enter the desired step size and size of the sliding window.

Results can be plotted using **Results** -> **Chart** or viewed in a table via **Results** -> **Table**.

6.2 Linkage Disequilibrium

This generates a linkage disequilibrium data set from SNP data.

NOTE: It is important to use only filtered data sets (apply **Filter** -> **Sites** first) when estimating linkage disequilibrium, as a raw alignment with numerous invariant bases will take a very long time and consume a large amount of memory to calculate.

😑 🔿 😑 🛛 Linkage Disequilibrium
Select LD type: Sliding Window \$
LD Window Size: 50
Sliding Window LD with 153375 comparisons.
How to treat heterozygous calls: Set to missing \$
Accumulate R2 Results
Run Close

Linkage disequilibrium between any set of polymorphisms can be estimated by clicking on a filtered set of polymorphisms and then using **Analysis** \Box **Link. Diseq.** At this time, *D'*, *r2* and *P*-values will be estimated. The current version calculates LD between haplotypes with known phase only (unphased diploid genotypes are not supported; see PowerMarker or Arlequin for genotype support).

D' is the standardized disequilibrium coefficient, a useful statistic for determining whether recombination or homoplasy has occurred between a pair of alleles.

 r^2 represents the correlation between alleles at two loci, which is informative for evaluating the resolution of association approaches.

D' and r2 can be calculated²¹ when only two alleles are present. If multiple alleles are present, a weighted average of D' or r2 is calculated between the two loci²². This weighted average is determined by calculating D' or r2 for all possible combinations of alleles, and then weighting them according to the allele's frequency. *Note: It is not entirely certain that this procedure fully accounts for allele number effects.*

P-values are determined by two methods. If only two alleles are present at both loci, then a two-sided Fisher's Exact test is calculated. *Note: Previous editions of TASSEL used a one-sided test, but TASSEL version 1.0.8 and later use a two-sided test.*

If more than two alleles are present, permutations are used to calculate the proportion of permuted gamete distributions that are less probable then the observed gamete distribution under the null hypothesis of independence²¹.

When calculating linkage disequilibrium, users have the option of employing "Rapid Permutations." If this

option is selected, the algorithm will compute either a fixed number of permutations or run until 10 permutations are found that are more significant than the observed P-value. While this slightly reduces P-values, it also saves a large amount of computational time. If an unbiased p-value is desired, then the user must unselect the **"Rapid Permutations"** check box.

"Full Matrix LD" calculates LD for every combination of sites in the alignment. "Sliding Window LD" calculates LD for sites within a window of sites surrounding the current site. The LD Window Size determines the width of the window on one side of the current site.

Linkage disequilibrium results can be plotted using **Results** -> **LD Plot** or viewed in a table via (**Results** -> **Table**).

6.3 Cladogram

$\Theta \cap \Theta$	Create Tree
	☑ Save distance matrix
	Clustering Method
	Neighbor Joining 🗧 🗧
R	un Close

This function generates a tree or cladogram data set.

TASSEL produces neighbor-joining trees using only simple parsimony substitution models.

To retrieve cladogram data, first select genotypic data from the Data Tree and then click **Analysis** -> **Cladogram**. The resulting tree data and the corresponding matrix will appear as separate data sets on the Data Tree.

Results can be plotted using **Results** -> Archaeopteryx Tree.

6.4 Kinship

This function generates a kinship matrix from a genotype. To do so, first highlight SNP data then click on the "Analysis/Kinship" submenu. The resulting dialog box will then provide the option to select "scaled IBS" or "pairwise IBS". Clicking "OK" generates a kinship matrix.

When a genotype file is selected and "pairwise IBS", each element i,j of the kinship matrix that is generated is equal to the proportion of the SNPs which are different between taxon i and taxon j. Distance is calculated for

each pair of taxa, ignoring any sites that have a missing value for one of the taxa. The distance matrix is converted to a similarity matrix by subtracting all values from 2 then scaling so that the minimum value in the matrix is 0 and the maximum value is 2. Kinship can be derived from a set of random SNP data (a minimum of several hundred SNPs spread over the whole genome is recommended). This ad-hoc rescaling method was implemented in an earlier version of TASSEL in order to provide a reasonable estimate of additive genetic variance, but tends to overestimate that value. Rescaling does not affect its use for correcting for population structure. It only affects the estimate of additive genetic variance and, consequently, heritability.

To provide a better estimate of addivitive genetic variance, an alternative method can be used by selecting "scaled IBS". This method (from Endelman and Jannink, 2012) codes genotypes as 2, 1, or 0, equal to the count of one of the alleles at that locus. It then replaces missing genotype values with the average genotypic score at that locus before estimating a relationship matrix. Other methods of imputing genotypes prior to calculating Kinship may provide a better result. For instance, rather than using this default treatment of missing values, using the numerical genotype method followed by imputation described in section 3.3 before running Kinship is a reasonable alternative. When using numerical genotypes, Kinship always applies the "scaled IBS" method.

Users may also load their own kinship data using **Data** \Box **Load**. Kinship matrices can be calculated using the SPAGeDi software package (<u>http://www.ulb.ac.be/sciences/ecoevol/spagedi.html</u>). Comparisons of methods for calculating kinship can be found in the literature (*e.g.* Stich et al. 2008).

6.5 GLM (General Linear Model)

This function performs association analysis using a least squares fixed effects linear model.

TASSEL utilizes a fixed effects linear model to test for association between segregating sites and phenotypes. The analysis optionally accounts for population structure using covariates that indicate degree of membership in underlying populations. A main effects only model is automatically built using all variables in the input data. A separate model is built and solved for each trait and marker combination. Any factors, covariates, reps or locations are included in every model as main effects. How the data is used must be defined either in the input data files or using the **Trait Filter** after the data has been imported but before it has been joined with a genotype.

General Linear Model (GLM) can be run using a numeric data set only or using numeric data joined to genotype data. If only numeric data is selected, best linear unbiased estimates (BLUEs or least square means) will be generated for the taxa for each trait. [Note: only factors and covariates intended to control field variation should be included at this stage. Population structure covariates which are intended to control for marker effects should only be included when markers are also in the analysis.] If numeric data with genotypes are analyzed, each trait by marker combination will be tested and two reports will be produced, one containing trait by marker F-tests and the other containing allele estimates.

To run GLM, select a data set and then click the GLM button. A dialog box will pop-up to allow the user to indicate that a permutation test should be run and to allow the number of permutations to be changed. The permutation test will be run using the method suggested by Anderson and Ter Braak (2003), which calculates the predicted and residual values of the reduced model (contained all terms except markers) then permutes the residuals and adds them to the predicted values. When the GLM options dialog is closed, the user is presented with a dialog allowing the output to be saved to a file rather than stored in memory and displayed by TASSEL. This option is useful when the output is expected to be very large and risks exceeding available RAM.

T Marker Tes	t		a l galend			1.18	free feetings			-			x
Trait	Marker	Locus	Locus_pos	marker_F	marker_p	markerR2	markerDF	markerMS	errorDF	errorMS	modelDF	modelMS	
EarDia	PZB00859.1	1	157104	1.663	0.199	0.007	1	23.522	223	14.144	3	31.92	21 🔺
EarDia	PZA01271.1	1	1947984	0.005	0.942	0	1	0.076	222	14.081	3	22.8	38 📟
EarDia	PZA03613.2	1	2914066	0.144	0.705	0.001	1	2.126	227	14.791	3	27.22	24
EarDia	PZA03613.1	1	2914171	0.014	0.905	0	1	0.208	228	14.66	3	24.48	37
EarDia	PZA03614.2	1	2915078	0.018	0.894	0	1	0.261	215	14.742	3	35.40)1
EarDia	PZA03614.1	1	2915242	2.663	0.104	0.012	1	39.907	213	14.984	3	38.96	57
EarDia	PZA00258.3	1	2973508	0.42	0.517	0.002	1	6.089	209	14.487	3	25.43	38
EarDia	PZA02962.13	1	3205252	0.374	0.541	0.002	1	5.374	217	14.357	3	25.18	33 🔻
•													F
	Print Export (CSV) Export (Tab)												

The following table shows an example of the Marker Test output as viewed with Results/Table:

In addition to displaying the F-statistics and p-values for the requested F-tests, the table also contains markerR2, mean squares (MS) and degrees of freedom (DF) for the marker effect, for the model (corrected for the mean), and for error. If taxa are replicated (across reps or environments), then the markers are tested using the taxa within marker mean square. If taxa are unreplicated, then the residual mean square is used. MarkerR2 is the marginal R-squared for the marker calculated as SS Marker (after fitting all other model terms) / SS Total, where SS stands for sum of squares. The following table shows an example of the Allele Estimates output as viewed with Results/Table:

Trait	Marker	Obs	Locus	Locus_pos	Allele	Estimate
EarDia	PZB00859.1	181	1	157104	С	0.804
EarDia	PZB00859.1	46	1	157104	A	0
EarDia	PZA01271.1	117	1	1947984	G	0.039
EarDia	PZA01271.1	109	1	1947984	С	0
FarNia	D7A03613 0	60	1	2014066	c	0.213
<u>ج</u>						4

For each marker and trait combination, each marker allele is listed along with the number of observations for taxa carrying that allele (Obs), the locus (usually chromosome) and locus position of that marker, the allele, and the estimate of the effect of that allele. Because of the way that GLM codes alleles, the last allele estimate for a marker is always zero and the other allele estimates are relative to that.

6.6 MLM (Mixed Linear Model)

This conducts association analysis via a mixed linear model (MLM).

A mixed model is one which includes both fixed and random effects. Including random effects gives MLM the ability to incorporate information about relationships among individuals. When a genetic marker based kinship matrix (K) is used jointly with population structure (Q), the "Q+K" approach improves statistical power compared to "Q" only⁹. MLM can be described in Henderson's matrix notation²³ as follows:

 $\mathbf{y} = \mathbf{X}\boldsymbol{\beta} + \mathbf{Z}\mathbf{u} + \mathbf{e}$

where y is the vector of observations; β is an unknown vector containing fixed effects, including genetic marker and population structure (Q); u is an unknown vector of random additive genetic effects from multiple background QTL for individuals/lines; X and Z are the known design matrices; and e is the unobserved vector of random residual. The **u** and **e** vectors are assumed to be normally distributed with null mean and variance of

where $\mathbf{G} = {}^{\acute{0} \frac{2}{a}} \mathbf{K}$ with ${}^{\acute{0} \frac{2}{a}}$ as the additive genetic variance and \mathbf{K} as the kinship matrix. Homogeneous variance is assumed for the residual effect which means $\mathbf{R} = \mathbf{I} {}^{\acute{0} \frac{2}{e}}$, where ${}^{\acute{0} \frac{2}{e}}$ is the residual variance. The proportion of genetic variance over the total variance is defined as heritability (h²).



When K is derived from pedigrees, the elements of K equal 2*Probability(IBD), where IBD means that two alleles drawn at random are identical by descent. Generally, K calculated from markers is an IBS matrix. The resulting multiplier is then not σ_a^2 but some unknown constant times σ_a^2 . Some methods for calculating K, such as those implemented in SPaGEDI, actually use markers to develop an estimate of the IBD relationship matrix. For those values of K, the resulting variance estimate can be considered an estimate of σ_a^2 as long as the assumptions of the method used to derive K are not violated for the population being analyzed. One implication is that two different K matrices may give very different estimates of σ_a and heritability yet produce the same model fit and test of marker association.

TASSEL implements several methods to improve statistical power and reduce computing time. The Restricted Maximum Likelihood (REML) estimates of \dot{b}_a^2 and \dot{b}_e^2 are obtained through the Efficient Mixed-Model Association (EMMA) algorithm²⁴ which is much faster than the expectation and maximization (EM) algorithm²⁵.

TASSEL also implements a method called compression which reduces the dimensionality of the kinship matrix to reduce computational time and improve model fitting. When MLM is used without compression (compression = 1), each taxon belongs to its own group. At the other extreme, GLM can be interpreted as maximum compression (compression = n) with all taxa in a single group. In that case, it is not possible to estimate the random effect independently of error and $6^{\frac{2}{a}}$ is absorbed into $6^{\frac{2}{e}}$. Between these two extremes, taxa can be grouped using cluster analysis based on kinship. When n individuals are compressed into s clusters (groups), the kinship among individuals is replaced with the kinship among groups. At some grouping levels, dependent on the trait and population being analyzed, this compressed MLM has improved statistical power compared to the regular MLM⁴. The optimum grouping with the best model fit for MLM without fitting genetic markers has the best statistical power for an association test of markers⁴. TASSEL allows users to specify the compression level (average number of individuals per group), or to have the program determine the optimum grouping.

Similar to GLM, MLM performs an association test for each combination of traits and markers. TASSEL provides users several options: 1) to estimate genetic and residual variance for each combination; 2) to get these estimates once for each trait without fitting genetic markers and then to use those estimates to test markers; 3) to use a prior heritability estimate provided by the user. The second option, named P3D (population parameters previously determined), has the same statistical power as the first option⁴. Using the P3D method or using a prior heritability can be much faster than calculating heritability for each marker.

Using MLM is very similar to using GLM. The difference is that in addition to choosing the joint data set (or numerical data set), kinship data must also be highlighted before clicking the MLM button to show the MLM option dialog. The option of "No Compression" is the regular MLM which is equivalent to "Custom level=1". For data sets with large numbers of taxa, the optimal compression option may be considerably slower than no compression or user supplied compression. This is because the algorithm solves the model once for each of a series of compression levels in order to determine the optimal one.

All MLM analyses create two output tables, model statistics and model effects. If compression is used, the analysis creates three tables.

Trait	Marker	Locus	Site	df	F	p	errordf	markerR2	Genetic Var	Residual Var	-2LnLikelihood
dpoll	None			0			257		8.068	14.585	1,477.183
dpoll	PZB00859.1	1	157104	1	0.001	0.979	250	0	8.068	14.585	1,477.183
dpoll	PZA01271.1	1	1947984	1	4.339	0.038	248	0.015	8.068	14.585	1,477.183
dpoll	PZA03613.2	1	2914066	1	0.132	0.716	255	0	8.068	14.585	1,477.183
dpoll	PZA03613.1	1	2914171	1	2.829	0.094	256	0.01	8.068	14.585	1,477.183
dpoll	PZA03614.2	1	2915078	1	0.044	0.834	243	0	8.068	14.585	1,477.183
dpoll	PZA03614.1	1	2915242	1	0.788	0.375	241	0.003	8.068	14.585	1,477.183
dpoll	PZA00258.3	1	2973508	1	0.732	0.393	240	0.003	8.068	14.585	1,477.183
dpoll	PZA02962.13	1	3205252	1	0.967	0.326	244	0.004	8.068	14.585	1,477.183
dnoll	P7402962 14	1	3205262	1	0.026	0.873	239	n 1	8 068	14 585	1 477 183
•											•

statistics table shows the results of the tests for each trait. The first line is for the model with no markers. Following that is a single line for each marker tested. The columns labeled "df", "F", and "p" are the degrees of freedom, F, and p-value from the F distribution for the test of the marker. The column "errordf" is the degrees of freedom used for the denominator of the F-test. The column labeled "markerR2" is the R2 for the marker calculated based on a formula for R2 for a generalized least squares GLS) model as shown here.

The

The columns "Genetic Var", Residual Var", and "-2LnLikelihood" list $\sigma 2a$, $\sigma 2e$, and minus two times the model likelihood, respectively. When the P3D option is used, all of the values are the same for a given trait because they are only calculated once. A second table lists the estimated effects of each allele for each marker similar to the output for GLM. The compression results table shown below shows the likelihood, genetic variance, and error variance for each compression level tested during the optimization process. The meaning of groups and compression is discussed above in the description of the compression method. The compression level with the lowest value of -2LnLk is used for testing markers.

Trait	# groups	Compression	-2LnLk	Var_genetic	Var_error
dpoll	259	1	1,480.402	7.362	8.146
dpoll	248	1.044	1,479.47	7.635	7.81
dpoll	243	1.066	1,479.505	7.656	7.855
dpoll	238	1.088	1,481.049	7.338	8.416
dpoll	234	1.107	1,482.935	6.957	9.069
dpoll	229	1.131	1,483.301	6.904	9.261
dpoll	224	1.156	1,482.597	6.866	9.394
dpoll	220	1.177	1,486.718	6.172	10.576
dpoll	215	1.205	1,485.526	6.407	10.342
dpoll	211	1.227	1,486.045	6.21	10.709
dpoll	207	1.251	1,488.214	5.897	11.345
•					÷.

6.7 Genomic Selection (using Ridge Regression)

This function performs ridge regression to predict phenotypes from genotypes. It is one of the methods used for genomic selection (GS).

The input dataset must contain one or more phenotypes and numeric marker data. Optionally, it may also contain factors and covariates. The analysis is run by selecting the input dataset then clicking the "GS" button. Because no additional user input is needed, the analysis will run immediately after the button is clicked. All traits will be analyzed separately using all of the genotypes, factors, and covariates in the dataset. The output will consist of two new datasets for each trait. One of the datasets will contain genomic estimated breeding values (GEBVs) for each taxon and the other will contain BLUPs for each marker in the genotype file. The output datasets will appear in the "Numerical" folder, which holds the input data as well. The output datasets can in turn be used for subsequent analysis. For example, it could be joined with the input data so that the predicted values could be graphed against the original values.

Understanding the input data requirements is important to ensure that the results of the analysis will be correct and useful. Genotypes must be numeric with one column for each marker. It is expected that the markers are bi-allelic, with the homozygotes coded as 1 and -1 and the heterozygotes coded as 0. However, any reasonable coding scheme will work. For instance, missing data could be replaced by a probability resulting from imputation. If any genotype data is missing, it will be imputed as the average of the marker scores across all taxa for that marker. If a user prefers to use a different method of imputation, then the missing genotypes must be imputed before importing the data into TASSEL.

GEBVs will be calculated for all taxa in the dataset, including any lines that have missing phenotype data. A typical use of genomic selection is to predict GEBVs for a set of unphenotyped lines based on the performance of a training set. To do that a dataset containing both the genotypes to be predicted and the genotypes of the training set can be joined with a dataset containing the phenotypes of the training set using a union join. All taxa in the phenotype set should have genotypes. If an individual without genotype data is included, all the marker data for that individual will be imputed, which is not a generally useful thing to do.

6.8 Geno Summary



\varTheta 🔿 🔿 TASSEL (Trait Ar	nalysis by aSSociation, Evolution, and Linkage) 5.0.5
File Data Filter Analysis Results	Help
Data Data Sequence mdp_genotype Result Genotype Summary mdp_genotype_OverallSummary mdp_genotype_SiteSummary mdp_genotype_SiteSummary mdp_genotype_TaxaSummary mdp_genotype_TaxaSummary Mdp_genotype_TaxaSummary Mdp_genotype_TaxaSummary Mumber of columns: 2 Number of rows: 14 Number of elements: 28 Overall Summary of mdp_genotype	Stat Type Value Number of Taxa 281 Number of Sites 3093 Sites x Taxa 869133 Number Not Missing 837722 Proportion Not Missing 0.96386 Number Gametes 1.7383E6 Gametes Not Missing 0.96386 Gametes Not Missing 0.96386 Gametes Not Missing 0.96386 Gametes Not Missing 0.96386 Gametes Missing 0.96386 Gametes Missing 0.96386 Gametes Missing 0.96386 Gametes Missing 0.03614 Number Heterozygous 9622 Proportion Heterozygous 0.01107
class net.maizegenetics.util.SimpleTableRep	ort

- Number of Taxa Number of Taxa in data set.
- Number of Sites Number of Sites in data set.
- Sites x Taxa Number of sites multiplied by number of taxa.
- Number Not Missing Number allele values not unknown (NN)
- Proportion Not Missing Number Not Missing / Sites x Taxa
- Number Missing Number unknown (NN) values
- Proportion Missing Number Missing / Sites x Taxa

- Number Gametes Number of Sites x Number Taxa x 2
- Gametes Not Missing Number of gametes not unknown
- Proportion Gametes Not Missing Gametes Not Missing / Number Gametes
- Gametes Missing Number unknown (N) gametes
- Proportion Gametes Missing Gametes Missing / Number Gametes
- Number Heterozygous Number of heterozygous values
- Proportion Heterozygous Number Heterozygous / Sites x Taxa

O O TASSEL (Trait A	nalysis b	y aSSociatio	on, Eve	olution, and	Linkage) 5.0.5	
File Data Filter Analysis Results	Help					
🚔 Data	Alleles	Numbe	r	Proportion	Frequency	
		C 24	6327	0.28342	0.29404	
mdn. construe		G 23	5079	0.27048	0.28062	
map_genotype		T 17	8046	0.20485	0.21254	
Result		A 16	8648	0.19404	0.20132	
🔻 🚞 Genotype Summary		N 3	1411	0.03614	0.0375	
mdp_genotype_OverallSummary		Y	3938	0.00453	0.0047	
mdp genotype AlleleSummary		R	2698	0.0031	0.00322	
mdp. genotype. SiteSummary		S	994	0.00114	0.00119	
mdp_genotype_sitesummary		К	839	9.6533E-4	0.001	
mup_genotype_raxasummary		w	596	6.8574E-4	7.1145E-4	
		M	557	6.4087E-4	6.649E-4	
		C:T	547	0.17685	NaN	
		G:A	486	0.15713	NaN	
e		1:0	408	0.13191	NaN	
Table Title: Allele Summary		A:G	3/3	0.12059	NaN	
Number of columns: 4		G:C	219	0.07081	NaN	
Number of rows: 27		C:A	184	0.05949	NaN	
Number of elements: 108		GI	165	0.05335	NaN	
Allele Summary of mdp_genotype		C:G	122	0.05011	NaN	
		A: I T: A	121	0.03912	NaN	
		T:A	100	0.03330	NaN	
		1.0	100	0.03492	NaN	
		C.C	50	0.02522	NaN	
		C.C	46	0.01017	NaN	
		Δ-Δ	24	0.0176	NaN	
*		TT	10	0.00614	NaN	
		1.1	19	0.00014	ivaliv	
lass net maizegenetics util SimpleTableRen	ort					

- Alleles Allele values present in data set. Single letter values are diploid where some letter represent heterozygous. Two letter values are major / minor combinations with count of sites.
- Number Number of occurrences
- Proportion Percentage the value occurs in data set.
- Frequency Percentage the value occurs in data set not counting unknown (N) values.

O O TASSEL (Trait A	nalysis by aSS	ociation, Ev	olution, and	Linkage) 5.0	0.5		
File Data Filter Analysis Results	Help						
🗖 Data	Site Number	Site Name	Chromoso	Physical Po	Number of Major	Allele Majo	10
	0	PZB0085	1	157104	281	C	
	1	PZA0127	1	1947984	281	G	Ĩ
map_genotype	2	PZA0361	1	2914066	281	Т	
Result	3	PZA0361	1	2914171	281	Т	
🔻 🚞 Genotype Summary	4	PZA0361	1	2915078	281	G	
mdp_genotype_OverallSummary	5	PZA0361	1	2915242	281	Т	
mdp genotype AlleleSummary	6	PZA0025	1	2973508	281	C	
mdn. genotyne. SiteSummany	7	PZA0296	1	3205252	281	Т	
mup_genotype_sitesummary	8	PZA0296	1	3205262	281	С	
mdp_genotype_TaxaSummary	9	PZA0059	1	3206090	281	Т	
	10	PZA0212	1	3706018	281	C	
	11	PZA0039	1	4175293	281	Т	
	12	PZA0286	1	4429897	281	C	
	13	PZA0286	1	4429927	281	С	
Table Title: Site Summary	14	PZA0286	1	4430055	281	Т	
Number of columns: 35	15	PZA0203	1	4490461	281	A	
Number of rows: 3093	16	zagl1.5	1	4835434	281	A	
Number of elements: 108255	17	zagl1.2	1	4835558	281	C	
Site Summary of mdn. genotype	18	zagl1.6	1	4835658	281	Т	
site summary of map_genotype	19	PZD0008	1	4836542	281	C	
	20	zagl1.1	1	4912526	281	A	
	21	PZB0091	1	5353319	281	C	
	22	PZB0091	1	5353655	281	G	
	23	PHM2244	1	5562502	281	G	
	24	PZA0309	1	8075572	281	G	
	25	PZA0018	1	8366368	281	G	
	26	PZA0018	1	8366411	281	Т	
class net.maizegenetics.util.SimpleTableRep	ort						

- _____
- Site Number Index of site
- Site Name Name of site
- Chromosome Chromosome
- Physical Position Physical Position on Chromosome
- Number of Taxa Number of taxa for site (same of all)
- Major Allele The major allele of site
- Major Allele Gametes Number of times major allele occurs for site (up to twice number of taxa)
- Major Allele Proportion Major Allele Gametes / (Number of Taxa * 2). Number of Taxa * 2 is the Number of Gametes for a Site.
- Major Allele Frequency Major Allele Gametes / ((Number of Taxa * 2) Gametes Missing)
- Minor Allele The minor allele of site
- Minor Allele Gametes Number of times minor allele occurs for site
- Minor Allele Proportion Minor Allele Gametes / (Number of Taxa * 2). Number of Taxa * 2 is the Number of Gametes for a Site.
- Minor Allele Frequency Minor Allele Gametes / ((Number of Taxa * 2) Gametes Missing)
- Gametes Missing Number of gametes with unknown (N) value
- Proportion Missing Gametes Missing / (Number of Taxa * 2)
- Number Heterozygous Number of taxa that are heterozygous for site.
- Proportion Heterozygous Number Heterozygous / Number of Taxa (not counting taxa that are unknown (NN))
- Inbreeding Coefficient -

• Inbreeding Coefficient Scaled by Missing -

File Data Filter Analysis Results	Help					
Data	Taxa	Taxa Name Nu	mber of Gan	netes M P	roportion N	lumber He F
	0	33-16	3093	190	0.03071	33
Sequence	1	38-11	3093	78	0.01261	22
mdp_genotype	2	4226	3093	176	0.02845	27
📄 Result	3	4722	3093	790	0.12771	147
🔻 🚞 Genotype Summary	4	A188	3093	158	0.02554	25
mdp genotype OverallSummary	5	A214N	3093	118	0.01908	25
mdp. genotype. AlleleSummary	6	A239	3093	76	0.01229	31
and a senst us Site Summary	7	A272	3093	330	0.05335	50
mdp_genotype_sitesummary	8	A441-5	3093	80	0.01293	26
mdp_genotype_TaxaSummary	9	A554	3093	104	0.01681	34
	10	A556	3093	254	0.04106	25
	11	A6	3093	78	0.01261	36
	12	A619	3093	124	0.02005	38
	^ 13	A632	3093	98	0.01584	32
Table Title: Taxa Summany	14	A634	3093	114	0.01843	33
Number of columns: 0	15	A635	3093	150	0.02425	40
Number of cours: 281	16	A641	3093	142	0.02296	26
Number of elements: 2520	17	A654	3093	226	0.03653	31
Taxa Summany of mdn. genetyne	18	A659	3093	160	0.02586	31
raxa summary of mup_genotype	19	A661	3093	468	0.07565	29
	20	A679	3093	140	0.02263	29
	21	A680	3093	128	0.02069	44
	22	A682	3093	112	0.01811	33
	23	AB28A	3093	238	0.03847	25
	24	B10	3093	118	0.01908	36
	25	B103	3093	136	0.02199	29
· · · · · · · · · · · · · · · · · · ·	26	B104	3093	112	0.01811	30

- Taxa Index of taxa.
- Taxa Name Name of taxa
- Number of Sites Number of sites for taxon (same for all).
- Gametes Missing Number of gametes with unknown (N) value. Every taxa / site combination has two gametes.
- Proportion Missing Gametes Missing / (Number of Sites * 2)
- Number Heterozygous Number of sites that are heterozygous for taxon
- Proportion Heterozygous Number Heterozygous / Number of Sites (not counting sites that are unknown (NN))
- Inbreeding Coefficient -
- Inbreeding Coefficient Scaled by Missing -

6.9 Stepwise

7 Results Menu

Results consists of the functions to present data as table or graphics.

7.1 Table

Allows data to be displayed in a spreadsheet view and exported into a flat file.

To create a table, select a data set from the Data Tree panel, then click on the menu "Results -> Table".



Shown below is an example in which the Taxa Summary is displayed.

9	Taxa Name	Number of Sites	Gametes Missing	Proportion Mis	Number Heter	Proportion He	Inbreeding Co	Inbreeding Co
	0 33-16	3093	190	0.031	33	0.011	Inbreeding Coe	ICSBM
	1 38-11	3093	78	0.013	22	0.007	Inbreeding Coe	ICSBM
	2 4226	3093	176	0.028	27	0.009	Inbreeding Coe	ICSBM
	3 4722	3093	790	0.128	147	0.054	Inbreeding Coe	ICSBM
	4 A188	3093	158	0.026	25	0.008	Inbreeding Coe	ICSBM
	5 A214N	3093	118	0.019	25	0.008	Inbreeding Coe	ICSBM
	6 A239	3093	76	0.012	31	0.01	Inbreeding Coe	ICSBM
	7 A272	3093	330	0.053	50	0.017	Inbreeding Coe	ICSBM
	8 A441-5	3093	80	0.013	26	0.009	Inbreeding Coe	ICSBM
	9 A554	3093	104	0.017	34	0.011	Inbreeding Coe	ICSBM
	10 A556	3093	254	0.041	25	0.008	Inbreeding Coe	ICSBM
	11 A6	3093	78	0.013	36	0.012	Inbreeding Coe	ICSBM
	12 A619	3093	124	0.02	38	0.013	Inbreeding Coe	ICSBM
	13 A632	3093	98	0.016	32	0.011	Inbreeding Coe	ICSBM
	14 A634	3093	114	0.018	33	0.011	Inbreeding Coe	ICSBM
	15 A635	3093	150	0.024	40	0.013	Inbreeding Coe	ICSBM
	16 A641	3093	142	0.023	26	0.009	Inbreeding Coe	ICSBM
	17 A654	3093	226	0.037	31	0.01	Inbreeding Coe	ICSBM
	18 A659	3093	160	0.026	31	0.01	Inbreeding Coe	ICSBM
	19 A661	3093	468	0.076	29	0.01	Inbreeding Coe	ICSBM
	20 A679	3093	140	0.023	29	0.01	Inbreeding Coe	ICSBM
	21 A680	3093	128	0.021	44	0.015	Inbreeding Coe	ICSBM
	22 A682	3093	112	0.018	33	0.011	Inbreeding Coe	ICSBM
	23 AB28A	3093	238	0.038	25	0.008	Inbreeding Coe	ICSBM
	24 B10	3093	118	0.019	36	0.012	Inbreeding Coe	ICSBM
	25 B103	3093	136	0.022	29	0.01	Inbreeding Coe	ICSBM
	26 B104	3093	112	0.018	30	0.01	Inbreeding Coe	ICSBM
	27 B105	3093	80	0.013	29	0.009	Inbreeding Coe	ICSBM
	28 B109	3093	284	0.046	43	0.015	Inbreeding Coe	ICSBM
	29 B115	3093	114	0.018	26	0.009	Inbreeding Coe	ICSBM
	30 B14A	3093	68	0.011	36	0.012	Inbreeding Coe	ICSBM
	31 B164	3093	68	0.011	26	0.008	Inbreeding Coe	ICSBM
	32 B2	3093	54	0.009	34	0.011	Inbreeding Coe	ICSBM
	33 B37	3093	90	0.015	26	0.009	Inbreeding Coe	ICSBM
	34 B46	3093	244	0.039	30	0.01	Inbreeding Coe	ICSBM
	35 B52	3093	122	0.02	19	0.006	Inbreeding Coe	ICSBM
	36 B57	3093	116	0.019	17	0.006	Inbreeding Coe	ICSBM
	37 B64	3093	156	0.025	28	0.009	Inbreeding Coe	ICSBM
	38 B68	3093	144	0.023	33	0.011	Inbreeding Coe	ICSBM
	39 B73	3093	56	0.009	23	0.008	Inbreeding Coe	ICSBM
	10 0 20 00 04	2002	1.40	0.024		0.010	h-1	TOODM

Data can be sorted by clicking on the column header of interest. A secondary sort can be done by holding down the CTRL key and clicking on a second column.

Data can be exported to flat files that are either comma-separated (Comma Separated Values = CSV) or tab-delimited. Both these formats can then be imported into a spreadsheet program such as Excel. Tables can also be printed.

7.2 Archaeopteryx Tree

Select a "Tree:..." data set to use. https://sites.google.com/site/cmzmasek/home/software/archaeopteryx





7.3 2D Plot

Displays 2D plots and determines color thresholds.

This function is useful for plotting associations in multiple environments.

First, select the desired result set. Using the drop down boxes provided, populate rows with "Environment," columns with "Site," and value with "PermuteP." The cutoff value for coloring can be chosen either by inputting a value in the text box or by using the slider tool to the right of the text box. Users can "mouse over" any box to view the value associated with that box, as shown here:

T 2-D chart										
🔀 📇 🏝 Cell size 🛛 14 🔽 🗋 only upper triangle 🔲 P-Value 🏢										
	Row: 🔽	Environme	nt 🔽 Colum	n: 🗹 Site	~	Value: Perm	uteP 🔽			
Cutoff 0.001 Save SVG										
PermuteP	18	511	625	880	1000	1459	1490	1570	1618	
CLAYTON.ID15										
HOMESTEAD.ID1										
Statistics - Min: 0.0 Max: 0).98 Mean: 0	.2658788 SD:	0.32200772				HOME	STEAD.ID1:	1490(0.0)	

If P-value coloring is desired, simply check the P-value box as shown below:

T 2-D chart										
💌 📇 🗛 Cell size 🛛 14 🔽 🗋 only upper triangle 🔽 P-Value ?										
	Row: 🔽	Environme	nt 🔽 Columi	n: 🗹 Site	v	Value: Permu	iteP 🔽			
Cutoff 80.9										
PermuteP	18	511	625	880	1000	1459	1490	1570	1618	
CLAYTON.ID15 HOMESTEAD.ID1										
Statistics - Min: 0.0 Max: 0).98 Mean: 0.	2658788 SD:	0.32200772			HOM	ESTEAD.ID1:	1459(0.04)		

By checking the P-value box, Cutoff selection tools will be disabled and fields will instead be colored according to the following grayscale:

Dialog 🛛 🔀
< 0.001
0.001 - 0.01
0.01 - 0.05
> 0.05
OK Cancel

This key can be shown by clicking on the "?" icon next to the P-value check box.

7.4 LD Plot

Displays the results from a linkage disequilibrium analysis.

After selecting the desired result from the Data Tree, choose Results -> LD Plot

The graph that is generated displays LD between pairs of sites calculated with the analysis step. The black diagonal represents LD between each site and itself. The default setting graphs r^2 in the upper right and *p*-values

in the lower left. This default can be modified by clicking on the buttons in the lower left. The left side of the graph contains a text description with the Chromosome and the Site name. At the bottom of the graph is a display of the position of each site along the chromosome. This display can be hidden by deselecting the "Schematic" check box. Legends that describe the color scheme appear on the right hand side of the graph.

The number of sites displayed can be selected by entering a number in the white box in the upper right corner or by moving the sliding bar next to it. To move through the graph use the sliding bars on the right and bottom. The red box in the small white window in the upper left corner will show what portion of the graph is displayed. To move only around the diagonal select the "Lock Y Axis to X" check box (recommended when visualizing a LD by sliding window analysis).



LD plots can be saved in several formats. The Save button will save the area of the graph shown in the screen, while the Save All button will save the entire graph.

7.5 Chart

Chart provides a variety of graphs for visualizing numeric data.

This feature can be used to display histograms, XY plots, bar charts and/or pie charts. Any numeric table data can be charted, including LD results, phenotypic data, diversity results, and association results.

Histograms: Use the graph type combo box to select the desired graph type (Histogram) from the list of options. Up to two different series of data can be plotted together. Users may specify the number of bins to be used in the histogram.



Scatter plots: Use the graph type combo box to select the desired graph type (XY Plot) from the list of options. Select data to be plotted in X and Y axes using the appropriate drop down boxes. If two data series are plotted simultaneously on the Y axis, the "2 Y Axes" checkbox will provide an axis for each.



7.6 QQ Plot

7.7 Manhattan Plot

8 GBS Menu

http://www.maizegenetics.net/tassel/docs/TasselPipelineGBS.pdf

9 Help Menu

Help provides information Tassel and diagnostics.

9.1 Help Manual

- 9.2 About
- 9.3 Show Memory
- 9.4 Logging

10 Tutorial

This tutorial reviews several common scenarios for using TASSEL in order to help the user better understand its capabilities for data manipulation and association analyses. The TASSEL software package includes a tutorial data set that can be downloaded from the TASSEL website (please unzip all files to a directory of your choice). This tutorial data set contains data for phenotype, genotype, population structure, and kinship.

10.1 Missing Phenotype Imputation

The phenotype file **mdp_traits** will be used to demonstrate the process of imputing missing data. Note that the data set below contains missing values (NaN).

ATASSEL (Trait Analysis by aSSociation, Evolution, and Linkage) 3.0.37						X
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		811	59.5	NaN	NaN	
		33-16	64.75	64.5	NaN	
mdp_bopaddon_ordecare		38-11	92.25	68.5	37.897	
Matrix		4226	65.5	59.5	32.21933	
mdp kinship	-	4722	81.13	71.5	32.421	
	_	A188	27.5	62	31.419	
Table Title: Phenotypes		A214N	65	69	32.006	
Number of columns: 4	=	A239	47.88	61	36.064	
Number of rows: 301		A272	35.63	70	NaN	
Number of elements: 1204	-	A441-5	53.5	67.5	35.008	
Mander of Creatinop, 1201		A554	38.5	66	33.41775	
Lond Cancel	1	A556	28	65	31.929	-
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To impute missing data, first select the mdp_traits data set in the Data Tree Panel and then click the **Transform** button (**Data** \Box **Transform**). The "Transform Column Data" window will open. Click on the **Impute** tab in this window. Finally, click on the **Create Data set** button to create the new data set with missing values imputed.

Note that missing values are now filled.



10.2 Principal Component Analysis

Principal component analysis (PCA) is a statistical tool that transforms a set of correlated variables into a smaller number of uncorrelated variables called principal components (PCs). The first PC captures as much of the variation as possible, and the succeeding PCs account for a decreasing fraction of the remaining variance. Another application of PCA is to use PCs derived from genetic markers to represent population structure⁸. This method requires much less computing time than maximum likelihood estimation. As most marker data are characters, numericalization must be performed first. A common approach for converting character marker scores is to set one of the homozygotes to 0, the other homozygote to 2, and the heterozygote to 1. For haploids, the conversion can be simply performed by coding one allele as 0 and the other as 1. The TRANSFORM function in TASSEL converts the major allele to 0. All the other alleles are collapsed to a single class and coded as 1. PCA requires that all variables should have variation and should not have missing values. As a result, filtering genotype to eliminate monomorphic markers and imputing missing values may be necessary. Imputing missing values can be done before or after numericalization. Here we demonstrate how to generate PCs from the genotype file in the tutorial data.

- 1. Remove monomorphic sites: Make sure TASSEL is in **Data** mode. Highlight the genotype and click **Site**. Set the minimum frequency to 0.05 and have "Remove minor SNP status" checked. Click **Filter**.
- 2. Numericalization: Highlight the filtered genotype and click **Transform**. Use the default option of "Collapse non major alleles." Click **Create data set**.
- **3**. Imputation of missing values: Highlight the numerical genotype and click **Transform** and then click **Impute** Tab. Use the default options. Click **Create data set**.
- 4. PCA: Highlight the imputed numerical genotype, click **Transform**, and then click **PCA** Tab. Change the default option to "Components=3" by choosing **Components** and type 3 in the text box. Click **Create data set**.

Filter Alignment Minimum Count: Minimum Frequency: Position Type: Start Position:	Filter Alignment 210 0.05 Position index 0	out of 281 sequences Physical Position (AGP) 157104	Numerical Genotype
End Position: Extract Indels Remove minor SNP states Generate haplotypes via sliding window Haplotype Length	2560 of 2561 sites	148907116	 Collapse Non Major Alleles Separate Alleles
Skep Length	Filter Cancel		Create Dataset Close

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🖓 🦳 Sequence	33-16	0	1	1	0	0	0	1	0	0	1
mdp_genotype	38-11	0	0	1	0	0	0	0	0	0	0
• mdp_genotype	4226	0	1	1	0	0	0	0	0	0	1
mdp_genotype	4722	0	0	1	0	0	0	NaN	0	0	0
mdp_genotype_chr1_157104-148907116	A188	1	1	1	0	0	0	0	0	0	0
Polymorphisms	A214N	0	1	0	1	0	1	0	0	0	0
Numerical	A239	1	1	0	0	1	1	0	0	0	0
mdp_population_structure	A272	1	1	0	0	1	1	1	0	0	0
mdp_traits	A441-5	0	1	1	0	0	0	0	0	0	0
mdp_genotype_chr1_157104-148907116_	A554	0	0	0	0	1	0	0	0	0	0
Matrix	A556	0	1	1	0	0	0	NaN	NaN	NaN	0
mdp_kinship	A6	1	1	0	0	1	1	0	0	0	0
····• Iree	A619	0	0	1	0	0	0	0	1	1	0
Fusions	A632	0	1	0	1	0	1	0	0	0	0
Synonymizer	A634	0	1	0	1	0	1	0	0	0	0
A Decult	A635	0	1	0	1	0	1	0	0	0	0
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	A659	1	0	0	0	1	0	0	0	0	0
Number of elements: 719922	A661	0	0	0	0	1	NaN	0	0	0	0
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S14.null	9.6	
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S16.null	4.3	
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a 🔨	Taxa	50	51	52	53	54	55	S6	57	58	59	
Sequence	33-16	0	1	1	0	0	0	1	0	0	1	~
mdp_genotype	38-11	0	0	1	0	0	0	0	0	0	0	
mdp_genotype	4226	0	1	1	0	0	0	0	0	0	1	
mdp_genotype	4722	0	0	1	0	0	0	0	0	0	0	
mdp_genotype_chr1_157104-148907116	A188	1	1	1	0	0	0	0	0	0	0	
Polymorphisms	A214N	0	1	0	1	0	1	0	0	0	0	
Numerical	A239	1	1	0	0	1	1	0	0	0	0	_
mdp_population_structure	A272	1	1	0	0	1	1	1	0	0	0	
mdp_traits	A441-5	0	1	1	0	0	0	0	0	0	0	_
mdp_genotype_chr1_157104-148907116_Collas	A554	0	0	0	0	1	0	0	0	0	0	_
mdp_genotype_chr1_15/104-14890/116_Collas	A556	0	1	1	0	0	0	0	0	0	0	_
Matrix	A6	1	1	0	0	1	1	0	0	0	0	_
• mdp_kinship	A619	0	0	1	0	0	0	0	1	1	0	_
Iree	A632	0	1	0	1	0	1	0	0	0	0	_
	A634	0	1	0	1	0	1	0	0	0	0	_
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imputed Phenotypic values.	A641	1	1	0	0	1	0	0	0	0	0	
Taya with insufficient data: 7	A654	0	0	0	0	1	0	0	0	0	0	
K = 20.9 (whether $k = 100$	A659	1	0	0	0	1	0	0	0	0	0	
K - 50.0% Cucoii):	A661	0	0	0	0	1	0.33333	0	0	0	0	
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Three items will be added to the data tree after running PCA. The first are the PCs. The second are the eigenvalues. And, the last are the eigenvectors. Here we use the Chart Function in the Result mode to graph the first three PCs, the individual eigenvalue contributions (sometimes called a skree plot) and the cumulative eigenvalue contributions. The eigenvalues are of interest because they equal the variance explained by each of the PCs.



10.3 Estimation of Kinship using genetic markers

While PCs can be used to capture major population subdivisions, kinship can be used to capture more subtle relationships. This section shows how to create a kinship matrix based on the same SNP data used to calculate PC's.

- 1. Remove monomorphic sites: Highlight the genotype and choose **Filter/Sites** on the menu bar. Set the threshold on MAF to 0.05, check "Remove minor SNP status," then click **Filter**.
- 2. Estimate kinship: Highlight the filtered genotype and click **Analysis/Kinship.** Leave "Scaled IBS" selected in the "Choose Kinship Method" dialog and click **OK**. A kinship matrix will be added to the data tree under Matrix category.
- 3. Alternatively, impute missing genotype data first then create the kinship matrix using the imputed data. To impute missing data, highlight the filtered genotype, choose Data/Transform, leave "Collapse Non-Major Alleles" selected, and click "Create Dataset". A new data set with "_Collapse" appended will appear in the "Numerical" folder. Highlight the collapsed data set, choose Data/Transform, select the Impute tab, then click "Create Dataset". Highlight the resulting imputed data then choose Analysis/Kinship.



10.4 Association analysis using GLM

We use three files from the tutorial data set to perform association analysis using the **GLM**. The first file mdp_genotype.hmp.txt, a set of SNPs scored at 3093 sites on 281 maize inbred lines. The second one is the population structure of 282 maize inbred lines (mdp_population_structure.txt). The last one is phenotypes for three traits, for 282 maize inbred lines (mdp_traits.txt). The statistical model is:

Flowering time = Population structure + Marker effect + residual

- 1. Remove monomorphic and low coverage sites: Highlight the mdp_genotype and click **Filter/Sites** on the menu bar. Set "Minimum Frequency" to 0.05, "Maximum Frequency" to 1.0, and "Minimum Count" to 150. Click **Filter** to create a filtered genotype data set.
- 2. Trait selection: Highlight the phenotype and click the menu item **Filter/Traits**. Uncheck all the traits except flowering time (DPOLL). Make sure that the Type is set to Data. Click **OK** to create a filtered phenotype.
- **3**. Covariate selection: The population structure is presented as the proportion of each population. There are three populations represented as Q1, Q2, and Q3. They sum to 100%. This creates linear dependency if we use all of them as covariates. While GLM can handle that properly, it will cause MLM to complain and refuse to complete your analysis. We can eliminate the dependency by removing one of the Q variables. In this demonstration, we exclude the last one. Highlight mdp_population_structure and click **Filter/Traits**. Uncheck the last population (Q3). Make sure that the Type is set to Covariate. Then click **OK** to create a filtered population structure data.
- 4. Joining data: Highlight the three filtered data sets by holding the Control key while selecting the individual data sets. Then click the menu item **Data/Intersect Join** to create a combined data set.
- 5. Association analysis: Highlight the joint data set then click the menu item **Analysis/GLM** to perform association analysis. Two reports will be added to the data tree.

	\varTheta 🔿 🖸 🛛 Filta	er Alignment			
		Filter Alignment			
	Minimum Count:	150	out of 281 sequences		
	Minimum Frequency:	0.05			
	Maximum Frequency:	1.0			
	Position Type:	Position index	7		
	Start Position:	0	-		
	End Position:	of 3092 sites			
	Remove minor SNP states Generate haplotypes via sliding window Haplotype Length Step Length Filter Sele	of 3092 sites	Cancel		
	Filter	ct Chromosomes	Cancel		X
Filter Traits / Modify Trait Properties		Filter Traits / Modif	fy Trait Properties		
Trait Type	Discrete Include	Trait	Type Discrete	Include	
EarHT data		Q1	covariate		
EarDia data		Q2 Q3	covariate		
Evolution	Selected Joclude Selected		Evolude Coloctod	e Selected	
				- Serviced	
Ex	clude All		Exclude All Include	a All	
	OK Cancel		OK Cance		

One of the reports added to data tree is labeled "GLM_Marker_Test_" followed by the name of the joint data. In addition to the information for traits and markers, the data set contains the following statistics:

marker_F: F value from the F test on marker;

marker_p: P value from the F test on marker;

marker $\overline{R2}$: R^2 for the marker after fitting other model terms (population structure);

markerDF: Degree freedom of marker;

markerMS: Mean square of marker;

errorDF: Degree freedom of residual error;

errorMS: Mean square of residual error;

modelDF: Degree freedom of model;

modelMS: Mean square of model.

	Trait		Marker	Locus	L	ocus_pos	marker_F	marker_p	markerR2	markerDF	markerMS	errorDF	errorMS	mod
Synonymizer		dpoll	PZB0085		1	157104	0.33532	0.71543	0.0018	2	7.32118	251	21.83356	
Result		dpoll	PZA0127		1	1947984	5.98887	0.01509	0.01593	1	130.91719	247	21.86006	
Diversity		dpoll	PZA0361		1	2914066	0.44396	0.50582	0.00117	1	9.7473	254	21.95558	
SNP Assavs		dpoll	PZA0361		1	2914171	1.94335	0.14533	0.00993	2	42.34854	256	21.79146	
D ID		dpoll	PZA0361		1	2915078	0.18011	0.67166	4.9717E-4	1	3.98879	242	22.14663	
		dpoll	PZA0361		1	2915242	1.17459	0.27955	0.00313	1	24.76818	240	21.08668	
Association		dpoll	PZA0025		1	2973508	1.31685	0.26993	0.00725	ā	28.6036	237	21.72128	
GLM_marker_test_Filtered_mdp_traits + F		dpoll	PZA0296		1	3205252	2.98033	0.05264	0.01559	ĩ	59.84505	243	20.08003	
GLM allele estimates for Filtered_mdp_tra		dpoll	PZA0296		1	3205262	0.33803	0.56153	9.1575E-4	1	6.53992	235	19.34731	
The Variances		dpoll	PZA0059		1	3206090	0.70844	0.49339	0.00369	2	15.59899	253	22.01874	
Stepwise		dpoll	PZA0212		1	3706018	0.18465	0.66777	4.8205E-4	1	4.09165	254	22.15916	
	C	dpoll	PZA0039		1	4175293	0.01174	0.91382	3.1912E-5	1	0.2533	243	21.58512	
C		dpoll	PZA0286		1	4429897	2.57509	0.1098	0.00668	1	56.3929	254	21.89943	
ble Title: Marker Test		dpoll	PZA0286		1	4429927	3.39142	0.03529	0.0176	1	72.99552	239	21.52361	
umber of columns: 13		dpoll	PZA0286		1	4430055	3.14175	0.04505	0.01722	1	68.18523	232	21.70296	
umber of rows: 2559		dpoll	PZA0203		1	4490461	0.73384	0.39245	0.00191	1	16.19389	254	22.06733	
umber of elements: 33267		dpoll	PZB0091		1	5353319	1.69532	0.1941	0.00445	1	36.29275	249	21.40765	
sts of Marker_Phenotyne AssociationCLM. fix		dpoll	PHM2244		1	5562502	1.29978	0.27446	0.00693	2	28.44728	246	21.88623	
		dpoll	PZA0309		1	8075572	0.09464	0.90973	4.9411E-4	2	2.08047	252	21.98287	
C		dpoll	PZA0018		1	8366368	0.14162	0.86803	7.639E-4	2	3.12032	243	22.03351	
		dpoll	PZA0018		1	8366411	4.48832	0.01214	0.02238	2	95.08609	256	21.18523	
		dpoll	PZA0052		1	8367944	0.98318	0.32245	0.00277	1	21.694	231	22.06508	
		deall	D740017		1	0010007	6 57604	0.0100	0.01640		140 61025	252	21 20067	

Clicking "marker_p" will sort the table by P value. The smallest P value is 3.5963×10^{-6} . A reasonable significance threshold is 1.9×10^{-5} , which is 5% after Bonferroni multiple test correction (0.05/2559). The denominator in the Bonferroni correction is the total number of SNPs tested. The association was significant.

The other data added to the data tree is labeled "GLM_Allele_Estimates_" followed by the name of the joint data. For the most significant SNP (highlighted in the figure below), there were two genotypes (AA and GG). There are 220 lines with genotype AA and 41 lines with allele GG. For the trait dpoll (days to pollination), the difference between the two homozygotes was 3.86 days.

	Trait Marker	Obs	Locus	Locus_pos Allele	Estimate	
Synonymizer	dpoll	PHM448.23	2	8 133775120	R 0	
Result	dpoll	PZA00766.1	133	8 133775220	T 4.06105	
Diversity	dpoll	PZA00766.1	115	8 133775220	C 2.47046	
SNP Assavs	dpoll	PZA00766.1	2	8 133775220	Y 0	
	dpoll	PZB01389.1	122	8 134723842	C -4.9284E	
	dpoll	PZB01389.1	137	8 134723842	т 0	
Association	dpoll	PZA03591.1	220	8 134813437	A 3.85777	
GLM_marker_test_Filtered_mdp_traits + F	c dpoll	PZA03591.1	41	8 134813437	G 0	
GLM allele estimates for Filtered_mdp_tra	dpoll	PZA03591.3	223	8 134813550	C 0.77616	
Variances	dpoll	PZA03591.3	33	8 134813550	т 0	
Stepwise	dpoll	PZA03591.2	104	8 134813696	G 1.06301	
Stebuise	dpoll	PZA03591.2	145	8 134813696	A 0	
0	dpoll	PZA00090.1	29	8 137480768	R 0.01513	
silmher of elements' 44499	dpoll	PZA00090.1	217	8 137480768	G 0.29198	
0	dpoll	PZA00090.1	14	8 137480768	A 0	
	lloqb	PZB00665.1	183	8 137572174	C 2.2406	

10.5 Association analysis using MLM

Running MLM in tassel is similar to running GLM. The difference is that in addition to the joint data (or numerical data), MLM requires kinship data to define the relationship between individuals. The kinship matrix

times a parameter equals the covariance matrix between individuals. Here we use kinship file from the tutorial data set to fit the following statistical model.

Flowering time = Population structure + Marker effect + Individuals + residual

Individuals and the residual are fit as random effects. The other terms are treated as fixed effects.

With respect to the marker effect, we will demonstrate the analysis using two sets of markers. One is the dwarf8 gene sequence used in the GLM tutorial. The other is a set of 3093 SNPs spread across the maize genome.

For the dwarf8 gene sequence, use the joint data set created by following the tutorial for GLM. Solve the mixed linear model by highlighting the joint data set and the kinship data then clicking the **MLM** button in **Analysis** mode.

ACCENT STATES AND A STATES AND	, and	l Linkage)	3.0.38	_	_		
File Tools Help <u>G</u> DPC							
ecces Data Analysis]	Delete	Wizard		Show Memory]	0%
🕂 Diversity	.ink.	Diseq.	🕅 Cladogram	AGTOG SNP Ex	tract 📑 Kin	ship 👫 GLM	
Data	*	Taxa	dpoll	Q1	Q2	Haplotype	
E Sequence		38-11	68.5	3.0E-3	0.993	NNC-CGCAT	A
 d8_sequence 		A272	70	1.9E-2	0.122	CTGTGATGC	
mdp_genotype		A441-5	67.5	5.0E-3	0.531	CTGTGATGC	
mdp_genotype	-	A554	66	1.9E-2	0.979	CTGTGATGC	=
mdp_genotype	-	A6	80.5	3.0E-3	3.0E-2	CTGTGATGC	
a8_sequence_cnr_6-2404		A619	61	9.0E-3	0.99	GCGCGACA	
Filtered_mop_traits + Filtered_mop_populat	•	A632	61	0.993	4.0E-3	CTGTGATGC	
Polymorphisms		B103	57.5	0.163	0.829	CTGTGATGC	
		B104	64.5	0.694	0.305	CTGTGATGC	
mdp_population_structure		B14A	63.5	0.998	1.0E-3	CTGTGATGC	
map_traits		B37	65.5	0.997	2.0E-3	CTGTGATGC	
Filtered_mdp_population_structure		B68	71.5	0.998	1.0E-3	CTGTGATGC	
Filtered_mop_traits		B73	70	0.999	1.0E-3	CTGTGATGC	
		B84	67.5	0.996	3.0E-3	CTGTGATGC	
		B97	60.5	1.6E-2	0.981	CTGTGATGC	
■ Iree	-	C103	66.5	2.0E-3	0.997	NNNNNNN	
		CI 187-2	70.5	6.0E-3	0.992	CTGTGATGC	
	-	CM105	65.5	0.995	4.0E-3	NNNNGACG	
• III • I		CM174	64	0.997	2.0E-3	GCGCGACA	
	· ·	CM7	61.5	5.9E-2	0.939	GCGCGACA	
< · · · · · · · · · · · · · · · · · · ·		CML 10	77	2.0E-3	2.0E-3	CTGTGATGC	_
Program Status	_	CM 247	20	2.05.2	15 OF 2	CTOTOATOC	

MLM Options
Compression Level
Optimum Level
Custom Level:
No Compression
Variance Component Estimation
P3D (estimate once)
Re-estimate after each marker
Run Cancel Help Me Choose

An MLM option dialog will pop up as shown above. Choose the default options, which use P3D and compression at the optimum compression level. After the Run button is clicked, the progress bar will start moving. The time required will depend on sample size, number of traits, number of markers, and the options chosen in the MLM option dialog. After the progress bar is reset to zero, indicating completion of MLM, three reports will be added to the data tree. The first two are similar to the reports created by GLM. The most significant SNP is still the

same, however the strength of association is weaker, with a P value of 7.199×10^{-4} (vs. 1.1021×10^{-4} from GLM) which does not pass the Bonferroni multiple test threshold (5×10^{-4}).

The third report contains the MLM specific statistics, including -2 Log Likelihood, genetic variance and residual variance components under different level of compression. These statistics are illustrated by the Chart function on the Result mode as follows.



In the example, 79 are included in the final analysis. When they are clustered into 44 groups, the -2 Log Likelihood reaches a minimum, which indicates the best model fit. The screening of SNPs was performed at this optimum compression level.

Note: When two or more individuals are clustered into one group, the variance component for the random effect is not equivalent to the one without compression. Consequently, the heritability derived should not be interpreted as the individual based heritability.

To perform a Genome-Wide Association Study (GWAS) on the 3093 SNPs, we need to create a new joint data set containing the filtered phenotype, population structure, and the genome-wide genotype. Highlight the new joint file and the kinship data and click the **MLM** button. Choose the default options on the MLM option dialog. The analysis will take a minute or two. The output report labeled "MLM_compression" indicates that 259 lines were used in the analysis. With 74 groups, the statistics from the best are as graphed below.



The strongest associated SNP is at 193565357 bp on chromosome 3. The P value is 1.3027×10^4 . The threshold is 3.2331×10^{-5} at significant level of 1% after Bonferroni multiple test correction (0.01/3093). The association was not significant. As illustrated below, the output labeled "GLM_Allele_Estimates" shows the marker effects assigned to genotypes for each SNP (The GLM is also the same). For example, the first SNP at 157104 bp on chromosome 1 had three genotypes (AA, CC and AC) coded as A, C, and M based on the IUPAC code, see Appendix (Nucleotide Codes).

ASSEL (Trait Analysis by aSSociation,	Evolution, and	d Linkage) 3.0.3	9			in the second se	_		
File Tools Help GDPC									
CARANG COCANG Data	Results	Delete	Wizard	M	Show Memory]	E	0%	
		Table 🕎 Tr	ee Plot	2D Plot	LD Plot	Chart			
Tree 🔺	Trait	Marker	Locus	Site	Allele	Effect	Obs		
+ Fusions	dpoll	PZB00859.1	1	157104	C	3.64912	197		
Synonymizer	dpoll	PZB00859.1	1	157104	A	3.60484	53		
📕 Result	dpoll	PZB00859.1	1	157104	M	0	3		
Oiversity	dpoll	PZA01271.1	1	1947984	С	-1.2325E0	121		
SNP Assays	dpoll	PZA01271.1	1	1947984	G	0	127		
• D	dpoll	PZA03613.2	1	2914066	G	0.22634	75		
🖃 🚽 📕 Association	dpoll	PZA03613.2	1	2914066	т	0	180		
MLM_statistics_for_Filtered_n	dpoll	PZA03613.1	1	2914171	т	5.20917	195		
MLM_compression_for_Filtere	dpoll	PZA03613.1	1	2914171	A	6.46891	61		
MLM_statistics_for_Filtered_n ≡	dpoll	PZA03613.1	1	2914171	w	0	2		
MLM_compression_for_Filtere	dpoll	PZA03614.2	1	2915078	G	-1.2702E-1	125		
MLM_statistics_for_Filtered_n	dpoll	PZA03614.2	1	2915078	A	0	118		
MLM_effects_for_Filtered_md	dpoll	PZA03614.1	1	2915242	т	0.55196	130		
MLM compression for Filtere	dpoll	PZA03614.1	1	2915242	A	0	111		
	dpoll	PZA00258.3	1	2973508	G	-2.7856E0	65		
Table Title: MLM effects ^	dpoll	PZA00258.3	1	2973508	с	-3.3585E0	175		
Number of columns: 7	dpoll	PZA00258.3	1	2973508	S	0	2		
	dpoll	PZA02962.13	1	3205252	т	-4.1007E0	218		
	dpoll	PZA02962.13	1	3205252	A	-3.2237E0	26		
	dpoll	PZA02962.13	1	3205252	w	0	3		
Load 📻	dpoll	PZA02962.14	1	3205262	С	-2.0366E-1	224		
	dpoll	PZA02962.14	1	3205262	G	0	15		
Export	dpoll	PZA00599.25	1	3206090	С	0.61187	26		
-	dpoll	PZA00599.25	1	3206090	т	-1.2877E-1	231		
Transform 💻	dpoll	PZA00599.25	1	3206090	Y	0	1		
	dpoll	PZA02129.1	1	3706018	т	0.59304	124		
	dpoll	PZA02129.1	1	3706018	С	0	131	· ·	
Program Status									

11 Appendix

Code	Meaning	
Α	A:A	
С	C:C	
G	G:G	
Т	T:T	
R	A:G	
Y	C:T	
S	C:G	
W	A:T	
К	G:T	
Μ	A:C	
+	+:+ (insertion homozygous)	
0	+:-	
-	-:- (deletion homozygous)	
Ν	Unknown	

11.2TASSEL Tutorial Data sets

http://www.maizegenetics.net/tassel/docs/TASSELTutorialData3.zip

File name	Туре	Format	
d8_sequence.phy	Genotype	Phylip Alignment	
mdp_genotype.hmp.txt	Genotype	Hapmap Alignment	
mdp_genotype.plk.ped	Genotype	Plink Alignment	
mdp_genotype.plk.map			
mdp_kinship.txt	Kinship	Numerical square matrix	
mdp_population_structure.txt	Population structure	Numerical trait data	
mdp_traits.txt	Phenotype	Numerical trait data	

File #1 is the sequence of dwarf8 gene with 2466 sites on 91 maize inbred lines. The data was described by the paper on the association between Dwarf8 and flowering time²⁶.

File #2-6 are 3093 SNPs on 281 maize association inbred lines. The data was presented in three formats (Hapmap, Plink and Flapjack). The data was created by the PANZEA project funded by NSF. Details of the data can be found at <u>http://www.panzea.org</u>.

File #5 and 6 are in pair for the format of Plink.

File #7 is kinship created by Yu et al.⁹.

File #8 is population structure of 282 maize inbred line²⁷.

File #9 is phenotype on three traits, including flowering time, on 282 maize inbred lines⁹.

11.3Frequently Asked Questions

1. What do I do if TASSEL misbehaves?

TASSEL is an open source software project hosted on SourceForge and has a bug tracking list at http://sf.net/projects/tassel where you can notify the developer community of problems. In order for a bug to be fixed, we must be able to replicate the problem. Thus, it is important to document the steps that were taken that produced the error. If the data you are working with is not too sensitive, please include the files which were used in the faulty procedure. If you would rather not post your data file on SourceForge, you may email it to one of the software developers.

2. Where do I turn for more information?

If you are having difficulty with a certain aspect of TASSEL, you can either email one of the software developers listed at <u>www.maizegenetics.net</u> or you may check the TASSEL forum on SourceForge <u>http://sf.net/projects/tassel</u>), as another user may have already addressed a similar question. There is also a TASSEL discussion group at <u>http://groups.google.com/group/tassel</u>.

3. How do I join the fun: TASSEL on SourceForge?

TASSEL is an open source project distributed under the GNU general public license. This means that the source code is available and the user is free to modify the code to suit their particular needs. We welcome input from developers and those who wish to become involved in the improvement of this software. The project is hosted on SourceForge (http://sf.net/projects/tassel), thereby allowing anyone to access the most recent changes to the code. This setup makes it convenient for anyone to add special functionality to TASSEL if they so desire. It also serves as a good platform for anyone who wishes to become involved in a bioinformatics software development project.

4. When I click on the most current version of TASSEL web start, a previous version appears. What should I do?

The previous version of TASSEL web start was cached in your machine. To replace it with the most current version, click the Start button in Windows, followed by Run. Type **javaws** and then click OK. In the window that opens, keep the most current version of TASSEL and delete the rest.

5. What should I substitute for missing values in TASSEL?

For numerical data in version 3 format, use NA or NaN. For numerical data in version 2 format, use "-999" for missing values. For SNP data, use "N". Kinship does not allow missing values.

6. Is it possible to change data names in the Data Tree?

Yes. Click on the desired data name in the Data Tree, wait for one second, and then click it again or immediately hit the F2 key. Rename the data set and then hit Enter to save the change.

7. How can I create a TASSEL icon on desktop?

Click "Start" on Microsoft Windows and select "Control Panel", then double click Java to show "java Control Panel". In "Temporary Internet Files" section, click "View" button show "Java Cache Viewer". Move mouse over TASSEL application and click right button and select "Install Shortcuts".

8. Why do I get empty squares in MLM association analysis?

The empty square means null information. The major reasons include non-convergence in the estimation of

variance components or that the statistic in question was not calculated. For example, marker F, p, and R² are not calculated when no marker is included in the model.

9. Why should I exclude one column of the population structure?

For some methods of calculating population structure, such as the software STRUCTURE, the population proportions sum to one. This produces linear dependence between the population co-variates. While the algorithm used by GLM tolerates that dependency, MLM will fail because the design matrix will not be invertible Excluding one column eliminates linear dependence between columns. Using PC axes to represent population structure does not result in linear dependency because all PC columns are guaranteed to be independent.

10. Can kinship replace population structure?

Sometimes. For some traits and populations, the K-only model may be as good as or better than the Q+K model. For others, Q+K may be superior. The Q-only model is not as effective for controlling population structure as the alternatives. Unfortunately, no general guidelines exist for predicting which model will perform best. As a result, an investigator may wish to fit all three models and compare the results. If eliminating false positives is very important, then it may make sense to accept the most conservative model. However, if the objective is to identify candidates for further study and the cost of following up on a false lead is low, the most liberal model may be preferred.

11. Why do TASSEL and SPAGeDi give different kinship estimates?

First, many algorithms exist to calculate kinship and their estimates will differ from one another. Secondly, the algorithm in TASSEL treats each genotype as a haplotype. It is not recommended that TASSEL be used to generate a kinship matrix from heterozygous genotype. In the near future, the TASSEL kinship algorithm will be modified to handle heterozygous diploids.

12. Can I get Marker R square using SAS Proc Mixed or TASSEL MLM?

SAS Proc Mixed does not produce an R² statistic. MLM in TASSEL does. The user manual describes how it is calculated.

13. Does MLM find more associations than GLM?

Sometimes. MLM has higher statistical power than GLM and may detect more true associations.. When the tested genetic markers are confounded with kinship structure, GLM does not correct for that as effectively as MLM and may produce more false positives

14. Do I need multiple test correction for the p value from Tassel? Yes.

15. Can TASSEL handle diploid genotype data?

While TASSEL accepts most common sequence alignment formats which handle polyploid genotype data including haploid and diploid, some analyses are not appropriate for heterozygous data. GLM or MLM fit SNPs one at a time, treating each distinct genotype as a separate class. This has the effect of fitting an additive plus dominance model. Separating the two effects is under consideration. Because handling heterozygotes as a third marker class is not appropriate for kinship or LD those analyses should not be used for that type of data at the present time. Work to improve handling heterozygotes is ongoing.

16. How to cite TASSEL?

The paper that describes TASSEL¹ as a software package and the papers that introduce specific methods implemented in TASSEL should be cited as appropriate, such as the unified ("Q+K") approach, EMMA, compression of mixed linear model and P3D. For example,:

- A. Linkage disequilibrium (D', R² and P value) were calculated by TASSEL¹.
- B. Association analyses were performed with the mixed linear model approach⁹ implemented by TASSEL¹.
- **C.** GWAS was performed with the compressed mixed linear model approach^{4,9} carried by TASSEL¹ which also implemented the EMMA³ and P3D⁴ algorithms to reduce computing time.

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